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### (57) Abstract

The present invention relates to a monoclonal antibody, preferably with specificity for CD38, having CDRs of foreign origin and a recipient framework region having a sequence of human or primate origin, wherein the original amino acid residues in position 29 and/or 78 of the sequence of the recipient framework region of the heavy chain is replaced by a replacement amino acid residue that is the same or similar to that in the corresponding position of the sequence of the corresponding framework region of the heavy chain of the antibody from which the CDRs are derived. Method of preparation of said antibody. Pharmaceutical composition containing said antibody. Use of said antibody for the treatment of cancer and autoimmune diseases.

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## Humanized antibodies to CD38

The present invention relates to antibodies and in particular to humanised antibodies and their preparation.

Antibodies typically comprise two heavy chains linked together by disulphide bonds and two light chains. Each light chain is linked to a respective heavy chain by disulphide bonds. Each heavy chain has at one end a variable domain followed by a number of constant domains. Each light chain has a variable domain at one end and a constant domain at its other end. The light chain variable domain is aligned with the variable domain of the heavy chain. The light chain constant domain is aligned with the first constant domain of the heavy chain. The constant domains in the light and heavy chains are not involved directly in binding the antibody to antigen.

The variable domains of each pair of light and heavy 20 The variable chains form the antigen binding site. domains on the light and heavy chains have the same general structure and each domain comprises a framework four regions, whose sequences are relatively 25 conserved, connected by three complementarity determining regions (CDRs: CDRL1, CDRL2, CDRL3, CDRH1, CDRH2 and CDRH3). The four framework regions largely adopt a betasheet conformation and the CDRs form loops connecting, and in some cases forming part of, the beta-sheet structure. The CDRs are held together in close proximity 30 by the framework regions and, with the CDRs from the other domain, contribute to the formation of the antigen binding site. The four framework regions are therefore crucial in ensuring the correct positioning of the CDRs relative to each other and hence in antibody binding. 35

The importance of the interaction between the CDRs and the framework regions has become increasingly evident as more and more non-human antibodies have become humanised, such humanised antibodies comprising non-human CDRs within a human framework. Humanised antibodies, in contrast to non-human antibodies, say mouse or rat antibodies, elicit a negligible immune response when administered to a human.

10 The prior art discloses several ways of producing such humanised antibodies. Thus EP-A-0239400 describes splicing CDRs into a human framework. Briefly, the CDRs are derived from a non-human species such as a rat or mouse whilst the framework regions of the variable domains, and the constant domains, are derived from a human antibody. Specifically, a humanised anti-CD52 antibody is disclosed in EP-A-0328404.

EP-A-054951 describes another way of humanising an antibody by re-shaping a non-human antibody to make it more like a human antibody. Basically, it comprises taking a non-human variable domain, such as mouse or rat variable domain, and changing the residues in the framework region to correspond to residues of a human framework.

In both EP-A-0239400 and EP-A-054951 an altered antibody is produced in which the CDRs of the variable domain of the antibody are derived from a first non-human species and the framework regions and, if present, the or each constant domain of the antibody are derived from human.

In such humanised antibodies a number of residues of the human framework region appear to exert a critical influ nce on the affinity of antigen binding (for example

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Kettleborough et al, 1991, Prot. Eng. 4:773). Certain positions in the heavy chain framework regions, particular, seem to be important in the retention of antigen-binding activity in a variety of antibodies. A number of investigators have reported the importance of residues at positions 67, 69 and 71, within the heavy chain framework region. These residues form a beta-sheet in contact with the interior aspect of the CDRH2 loop: presumably mismatches at these positions distort the CDR shape. Also, residues at positions 91 for correct CDRH3 and 94 appear to be important conformation in many heavy chains (for example Tempest et al, Bio/Technology 9:266). Other positions likely to affect antigen-binding are residues 27, 30 and 94 in the heavy chain, and residue 49 and 71 in the light chain (numbering according to the Kabat system). Furthermore, in the heavy chain the importance of regions 66-73 and ... 27-30 has been recognised in the literature, with residues 66-73 lying in close contact with CDRH2. It has now been found that the residues 29 and 78 of the framework region occupy a pocket which lies close to and affects antigen binding and that undesirable effect can be obviated by using residues corresponding to those in the corresponding position of the framework region of the antibody from which the CDRs are derived.

Accordingly, the present invention is directed to a monoclonal antibody having donor CDRs of foreign origin and a recipient framework region having a sequence of human or primate origin, wherein the original amino acid residue in position 29 or 78 of the sequence of the recipient framework region of the heavy chain is replaced by a replacement amino acid that is the same or similar to that in the corresponding position of the sequence of

the corresponding framework region of the antibody from which the CDRs are derived. By "similar" is meant an amino acid of equivalent size preferably of equivalent size, hydrophobicity and charge.

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Typically, the original amino acid residues in positions 29 and/or 78 of the recipient framework region are larger than their corresponding residues in the framework region of the antibody from which the CDRs are derived. Examples of these larger residues include tyrosine, histidine, tryptophan and 2-phenylalanine. Examples of the smaller corresponding residues in the framework region of the antibody donating the CDRs include glycine, alanine, valine, serine and leucine. In accordance with the invention, the larger original residue in positions 29 and/or 78 of the recipient framework is replaced with a replacement amino acid residue that is either the same or similar to the corresponding smaller residue of the antibody which is donating the CDRs.

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Although it is preferable for the replacement amino acid residue to be the same as the corresponding residue of the antibody which is donating the CDRs it can also be a similar amino acid residue provided the character with respect to size and preferably also hydrophobicity and charge is essentially the same i.e. conserved. For example, if the residue of the antibody which is donating the CDRs has a valine in position 29 and/or 78, then instead of having a replacement amino acid residue in the recipient framework which is also valine, one could, for example, use alanine instead since alanine equivalent charge, size and hydrophobicity to valine and thus similar. The use of a similar amino acid in place of the exact same amino acid is a practice which is well in the art and known as conservative establish d

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substitution.

By way of example, in a mouse heavy chain framework, side chains of Leu-29 and Val-78 would pack together in a small pocket close to CDRH1 whilst in the corresponding human heavy chain framework, such as for example NEW, which otherwise bears close homology to the mouse framework, the analogous positions are occupied by two Phe residues. The large aromatic side-chains appear to be too bulky to pack in the same fashion as in the mouse 10 antibody and so alter the disposition of neighbouring surface residues resulting in a different conformation of CDRH1 in a humanised antibody. Substituting either ... Phe residue by the smaller murine residue partially 15 relieves this effect allowing antigen binding. Full affinity is generally restored by replacement of both residues. It is therefore preferred that amino acids in both positions 29 and 78 are replaced.

In accordance with the invention, the replacement amino acid residues fit into the pocket without causing distortion of, for example, the CDRH1 conformation.

preferably, the framework of the antibody heavy chain is homologous to the corresponding framework of the human antibody NEW (Saul et al, J. Biol. Chem. 253:585-597, 1978). The final residue of framework 1 in this case is suitably Ser or Thr, preferably Ser. This residue is at position 30 (Kabat et al, 1987). Preferably the framework of the antibody light chain is homologous to the variable domain framework of the protein REI (Epp et al, Eur. J. Biochem., 45:513-524, 1974).

Particular examples of murine heavy chains in which residues 29 and 78 pack together in a small pocket close

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to CDRH1 are those in Kabat groups IB and IIC.

By contrast, other examples of human heavy chains which have bulky residues in positions 29 and 78 in the framework region are LES-C, T52, Ab44, HIGI and NEW, as listed in Kabat.

Species other than the mouse that may have residues of a small size in positions 29 and 78 are for example, the rat, rabbit and hamster.

All amino acid residue positions referred to herein employ the Kabat numbering system.

- An antibody according to the invention may be produced by a method including the steps of:
  - (i) obtaining the sequence of a donor heavy chain;
- 20 (ii) selecting a recipient human or primate framework by best-fit homology method;
  - (iii) replacing the amino acid residue in position 29 or 78 of the sequence of the recipient framework region of the heavy chain by an amino acid that is the same or similar to that in the corresponding position of the sequence of the corresponding framework region of the antibody from which the CDRs are derived.

The antibody heavy chain may be co-expressed with a complementary antibody light chain. At least the framework regions of the variable domain and the or each constant domain of the complementary chain generally are derived from the primate or human recipient. Preferably

the CDRs of both chains are derived from the same selected antibody.

The antibody preferably has the structure of a natural antibody or a fragment thereof. The term antibody may 5 therefore comprise a complete antibody, (Fab'), a fragment, a Fab fragment, Fv fragment, Fd fragment, SFv, a light chain dimer or a heavy chain and derivatives thereof. The antibody may be an IgG such as an IgG1, IgG2, IgG3 or IgG4, IgM, IgA, IgE or IgD. Furthermore, 10 the antibody may comprise modifications of all classes e.g. IgG dimers, Fc mutants that no longer bind Fc receptors or mediate Clq binding (blocking antibodies). The antibody may also be a chimeric antibody of the type described in WO 86/01533) which comprises an antigen 15 binding region and a non-immunoglobulin region. The antigen binding region is an antibody light chain variable domain or heavy chain variable domain. 💥 Typically, the antigen binding region comprises both light and heavy chain variable domains. 20 immunoglobulin region is fused at its C-terminus to the antigen binding region. The non-immunoglobulin region is typically a non-immunoglobulin protein and may be an enzyme, a toxin or a protein having known binding specificity. The two regions of the chimeric antibody 25 may be connected via a cleavable linker sequence.

The invention is preferably employed to humanise an antibody, for example, an antibody of rat, rabbit, hamster or mouse origin. The framework regions and constant domains of the humanised antibody are therefore of human or primate origin whilst the CDRs of the light and/or heavy chain of the antibody are for example, rat or mouse CDRs. The antibody may be a human or primate IgG such as IgG1, IgG2, IgG3, IgG4; IgM; IgA; IgE or IgD

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in which the CDRs are of rat or mouse origin.

The antibody from which the donor CDRs are derived is typically an antibody of a selected specificity. In order to ensure that this specificity is retained, either the variable domain framework regions of the antibody are re-shaped to correspond to variable domain framework regions of a human or primate antibody or the CDRs are grafted onto the closest human or primate framework regions. Either way, the resulting antibody preferably comprises non-human CDRs and human or primate framework regions that are homologous with the corresponding framework regions of the antibody from which the CDRs are derived. Preferably there is a homology of at least 50% between the two variable domains.

There are four general steps to produce a humanised antibody. These are:

- 20 (1) determining the nucleotide and predicted amino acid sequence of the light and heavy chain variable domains of the antibody from which the CDRs are derived;
- 25 (2) deciding which human or primate antibody framework region to use;
  - (3) the actual grafting or re-shaping methodologies/ techniques; and
  - (4) the transfection and expression of the grafted or re-shaped antibody.

These four steps are explained below.

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# Step 1: Determining the nucleotide and predicted amino acid sequence of the antibody light and heavy chain variable domains

To humanise an antibody the amino acid sequence of the non-human antibody's (donor antibody's) heavy and light chain variable domains needs to be known. The sequence of the constant domains is irrelevant. The simplest method of determining an antibody's variable domain amino acid sequence is from cloned cDNA encoding the heavy and light chain variable domain.

There are two general methods for cloning a given antibody's heavy and light chain variable domain cDNAs:

(1) via a conventional cDNA library, or (2) via the polymerase chain reaction (PCR). Both of these methods are widely known. Given the nucleotide sequence of the cDNAs, it is a simple matter to translate this information into the predicted amino acid sequence of the antibody variable domains.

## Step 2: Designing the humanised antibody

There are several factors to consider in deciding which

human antibody (recipient antibody) sequence to use
during humanisation. The humanisation of light and heavy
chains are considered independently of one another, but
the reasoning is basically the same.

This selection process is based on the following rationale: A given antibody's antigen specificity and affinity is primarily determined by the amino acid sequence of the variable region CDRs. Variable domain framework residues have little or no direct contribution.

The primary function of the framework regions is to hold

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the CDRs in their proper spacial orientation to recognise the antigen. Thus the substitution of rodent CDRs into a human variable domain framework is most likely to result in retention of the correct spacial orientation if the human variable domain is highly homologous to the rodent variable domain from which the CDRs were derived. A human variable domain should preferably be chosen therefore that is highly homologous to the rodent variable domain(s).

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A suitable human antibody variable domain sequence can be selected as follows:

Using a computer program, search all available protein (and DNA) databases for those human 15 antibody variable domain sequences that are most homologous, for example, to the rodent antibody This can be domains. accomplished with a program called FASTA but other suitable programs are available. 20 output of the program is a list of sequences most homologous to the rodent antibody, the percent homology to each sequence, and an alignment of each sequence to the rodent sequence. done independently for both the heavy and light 25 chain variable domain sequences. The above easily accomplished more are customised sub-databases are first created that only include human immunoglobulin sequences. two benefits. First, the actual computational time is greatly reduced because analyses are restricted to only those sequences of interest rather than all the sequences in the databases. The s cond benefit is that, restricting analyses to only human immunoglobulin 35

sequences, the output will not be cluttered by the presence of rodent immunoglobulin sequences. There are far more rodent immunoglobulin sequences in databases than there are human.

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List the human antibody variable domain sequences that have the most overall homology to the rodent antibody variable domain (from above). Do not make a distinction between homology within the framework regions and CDRs. Consider the overall homology.

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Eliminate from consideration those human sequences that have CDRs that have a different length than those of the rodent CDRs. This rule does not apply to CDR 3, because the length of this CDR is normally quite variable. Also, there are sometimes no or very few human sequences that have the same CDR lengths as that of the rodent antibody. If this is the case, this rule can be loosened, and human sequences with one or more differences in CDR length can be allowed.

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(iv) From the remaining human variable domains, one is selected that is most homologous to that of the rodent.

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(v) The actual humanised antibody (the end result) should contain CDRs derived from the rodent antibody and a variable domain framework from the human antibody chosen above.

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(vi) Instead of re-shaping or grafting to produce a humanised antibody, it would also be possible to synthesise the entire variable domain from scratch once the amino-acids of the non-human variable domain has been determined and the most homologous human variable domain has been identified.

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(vii) If donor heavy chain has two small residues at positions 29 and 78, and recipient chain has large, typically aromatic, residues at one or both of these positions, then further analysis is required.

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(viii) This analysis may take the form of a sequence comparison between the CDRH1 of the donor chain and that of other antibodies. For example, a CDRH1 sequence of SYGVH has been shown to require small residues at positions 29 and 78 for complete activity, and it is to be expected that other antibodies with the same or similar CDRH1 sequence will also require residues at these positions.

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Alternatively, the analysis may take the form of detailed computer aided modelling of the CDRH1 region of the proposed humanised antibody using standard techniques (for example the ADM package from Oxford Molecular Ltd). If this analysis, for example, reveals that CDRH1 lies in close approximation to the packed side chains of residues 29 and 78, and that altering these residues from human to smaller residues changes the orientation or position of CDRH1, then such smaller residues should replace the human ones. An example of such a perturbation of CDRH1 is shown in Figures 5 and 6.

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## Step 3: Grafting and re-shaping

See EP-A-0239400 and EP-A-054951 for details.

## 5 Step 4: The transfection and expression of the altered antibody

Once the antibody has been humanised and residues 29 and/or 78 replaced, the cDNAs are linked to the appropriate DNA encoding light or heavy chain constant region, cloned into an expression vector, and transfected into mammalian cells. These steps can be carried out in routine fashion. A humanised antibody may therefore be prepared by a process comprising:

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- (a) preparing a first replicable expression vector including a suitable promoter operably linked to a DNA sequence which encodes at least a variable domain of an Ig heavy or light chain, the variable domain comprising framework regions from a human or primate antibody and CDRs comprising at least parts of the CDRs from a second antibody of different origin;
- (b) if necessary, preparing a second replicable expression vector including a suitable promoter operably linked to a DNA sequence which encodes at least the variable domain of a complementary Ig light or heavy chain respectively;

- (c) transforming a cell line with the first or both vectors; and
- (d) culturing said transformed cell line to produce 35 said altered antibody.

Preferably the DNA sequence in step (a) encodes both the variable domain and the or each constant domain of the antibody chain, the or each constant domain being derived from the human or primate antibody. The antibody can be recovered and The cell purified. line which is transformed to produce the altered antibody may be a Chinese Hamster Ovary (CHO) cell line or an immortalised mammalian cell line, which is advantageously of lymphoid origin, such as a myeloma, hybridoma, trioma, or quadroma cell line. The cell line may also comprise a normal lymphoid cell, such as a B-cell, which has been immortalised by transformation with a virus, such as the Epstein-Barr virus. Most preferably, the immortalised cell line is a myeloma cell line or a derivative thereof.

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Although the cell line used to produce the altered antibody is preferably a mammalian cell line, any other suitable cell line, such as a bacterial cell line or a yeast cell line, may alternatively be used. In particular it is envisaged that <u>E. coli-derived bacterial</u> strains could be used.

Some immortalised lymphoid cell lines, such as myeloma cell lines, in their normal state secrete isolated Ig light or heavy chains. If such a cell line is transformed with the vector prepared in step (a), it may not be necessary to carry out step (b) of the process, provided that the normally secreted complementary to the variable domain of the Ig chain encoded by the vector prepared in step (a). where the immortalised cell line does not secrete a complementary chain, it will be necessary to carry out step may be carried out (b). This by manipulating the v ctor produced in step (a) so that this vector encodes not only th variable domain of an altered antibody light or heavy chain, but also the complementary variable domain.

Alternatively, step (b) is carried out by preparing a second vector which is used to transform the immortalised cell line. This alternative leads to easier construct preparation, but may be less preferred than the first alternative in that it may not lead to as efficient production of antibody.

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Where the immortalised cell line secretes a complementary light or heavy chain, the transformed cell line may be produced for example by transforming a suitable bacterial cell with the vector and then fusing the bacterial cell with the immortalised cell line by spheroplast fusion. Alternatively, the DNA may be directly introduced into the immortalised cell line by electroporation or other suitable method.

The present process has been applied to obtain an 20 antibody against the CD38 surface antigen.

Briefly, a humanised anti-CD38 monoclonal antibody (termed h3S) was produced in the following fashion. cDNA 25 was obtained from hybridoma cells secreting the murine cDNA clones monoclonal anti-(human CD38) AT13/5. encoding the heavy and light chains of the mouse antibody were identified and sequenced (Sequences 1 and 2 attached in Figures 1 and 2). This information was then used to choose appropriate human frameworks to receive the CDR grafts by the best-fit homology method. This procedure identified the REI light chain and the NEW heavy chain as the optimal choices.

CDRs were grafted on to the human frameworks. 35

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addition, guided by published work (Riechman et al., 1988 Nature 332: 323 and Tempest et al., 1991, Bio/Technology 9:266), four framework changes were made at this stage at positions likely to affect antigen-binding: residues 27,30 and 94 in the heavy chain, and residue 49 in the light chain (numbering according to the Kabat system). The resulting humanised antibody was tested for CD38 binding, with negative results. Expression of the humanised light chain together with a chimeric heavy chain (murine VH, human CH) produced functional antibody, indicating that the humanisation of the light chain was adequate.

A further series of heavy chain framework changes were examined. In particular, the analysis identified a stretch of sequence from residue 66 to 73 which lies in close contact with CDRH2 and a pocket formed by the side chains of residues 29 and 78, lying close to CDRH1, as affecting antigen binding. As mentioned earlier on the importance of the regions 66-73 and 27-30 is recognised in the literature, though the role of residue 29 and 78 and the interaction between the side chains of residues 29 and 78 is not.

Although the invention is described with reference to an anti-CD38 antibody it is applicable to any antibody, whatever antigen it binds to. In particular any antibodies that bind the 40kD antigen (CO/17.1.A) as disclosed in J. Cell. Biol., 125 (2) 437-446, April 1994 and in Proc. Natl. Acad. Sci. 87, 3542-3546, May 1990, carcinoma antigens and antigens involved in autoimmune diseases. A specific example of an anti-40kD antibody is 323/A3.

35 Another example of an antibody is an anti-folate receptor

antibody as disclosed in A. Tomasetti et al, Federation of European Biochemical Societies Vol 317, 143-146, Feb 1993. A specific example of an anti-folate antibody is MOV18. Further examples of antibodies include anti-CEA, mucin, anti-20/200KD, anti-ganglioside, digoxin, anti-CD4 and anti-CD23.

In particular the anti-CD38 antibody has the nucleotide sequences for the heavy chain and light chain variable region as shown in Figures 3, 3a and 4. 10

According to another aspect of the present invention there is provided the use of antibody according to the In particular there is present invention in therapy. provided the use of antibodies according to the invention for the treatment of cancer and their associated metastases and for treatment of autoimmune diseases, in particular for the treatment of multiple myeloma, lymphoma and rheumatoid arthritis.

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The anti-CD38 antibody of the present invention can be used in the treatment of multiple myeloma.

is a transmembrane glycoprotein expressed by CD38 immature B lymphocytes, activated T and B lymphocytes, 25 and plasma cells. Antibodies to CD38 capable of causing cell lysis may be useful in the immunotherapy of tumours bearing this antigen, principally multiple myeloma and 50% of non-Hodgin's lymphomas. Additionally, anti-CD38 antibodies may be useful in the treatment of autoimmune diseases such as rheumatoid arthritis and myaethenia gravis, as they have the potential to suppress both the humoral and cellular effector arms of the immune system.

A CD38 antibody according to the present invention has 35

been demonstrated to be lytic for cells expressing CD38 on their surface. The humanised antibody has been shown to bind CD38 and compete with the parental antibody in CD38 binding.

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Multiple myeloma is a neoplasm characterised by an accumulation of a clone of plasma cells, frequently accompanied by the secretion of immunoglobulin chains. Bone marrow invasion by the tumour is associated with anaemia, hypogammaglobinaemia and granulocytopaenia with concomitant bacterial infections. An abnormal cytokine environment, principally raised IL6 levels, often results in increased osteoclasis leading to bone pain, fractures and hypercalcaemia. Renal failure is not uncommon in the context of high concentrations of myeloma immunoglobulin and hypercalcaemia.

A variety of therapeutic protocols have been tried over recent years with little impact on the overall prognosis for myeloma patients. Treatment with melphalan and prednisolone remains the standard therapy, as it was thirty years ago (Bergsagel, 1989). A response to chemotherapy is associated with the induction remission with median duration of about two years, but in all cases this is followed by eventual relapse and death (Alexanian and Dimopoulos, 1994 New England J. of Medicine Vol. 330: 484). More aggressive chemotherapy utilising multiple cytotoxic agents has yielded little additional benefit in terms of survival or duration of remission, though high-dose therapy followed autologous bone marrow transplant remains an area of active development.

Several workers have proposed immunotherapeutic 35 strategies to combat myeloma. Interleukin 6 has been

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suggested to be a major growth factor for myeloma cells and may function in either an autocrine or paracrine fashion. Based on such results, interventions aimed at disrupting the IL6 signalling system have been designed. Two murine monoclonal that neutralise IL6 suppressed the proliferation of myeloma cells in a patient with leukaemic variant of the disease, though the tumour relapsed after 60 days.

10 Administration of anti-IL6 receptor monoclonal antibody to SCID mice engrafted with cells from a human myeloma cell line suppressed tumour growth, though only if the antibody was administered one day after injection of the myeloma cells. Antibody given after five days of growth had no significant effect. A CDR-grafted form of this antibody has also been prepared for possible human therapeutic use.

In a similar vein, myeloma cells bearing high levels of IL6 receptor have also been targeted by chimeric cytotoxinx consisting of IL6 variants linked to a modified form of <u>Pseudomonas</u> exotoxin. Cell killing is seen in vitro though the applicability of this technique in the clinic remains to be seen.

Our preference is for a more physiological approach, targeting myeloma cells for killing by the host immune system. The surface antigen CD38 is strongly expressed by more than 90% of multiple myeloma cells, and its suitability as a target for lytic immunotherapy has been discussed (Stevenson et al, 1991 Blood, Vol. 77, 5: 1071-1079). The same report also demonstrated the competence of effector cells from myeloma patients for lysis of target cells coated with a chimeric anti-CD38.

The dosages of such antibodies will vary with the condition being treated and the recipient of the treatment, but will be in the range 1 to about 100 mg for an adult patient, preferably 1 - 10 mg, usually administered daily for a period between 1 and 30 days. A two part dosing regime may be preferable wherein 1 - 5 mg are administered for 5 - 10 days followed by 6 - 15 mg for a further 5 - 10 days.

Also included within the invention are formulation 10 containing a purified preparation of an anti-CD38. Such formulation preferably include, in addition to antibody, a physiologically acceptable diluent or carrier possibly in admixture with other agents such as other antibodies or antibiotic. Suitable carriers include but are not limited to physiological saline, phosphate buffered saline, phosphate buffered saline glucose and buffered saline. Alternatively, the antibody may be lyophilised (freeze-dried) 20 reconstituted for use when needed, by the addition of an aqueous buffered solution as described above. Routes of administration are routinely parenteral including intravenous, intramuscular, subcutaneous and intraperitoneal injection or delivery.

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The following Examples illustrate the invention. In the accompanying drawings:

- Figure 1 shows the nucleotide and predicted amino acid sequence of mouse anti-CD38 antibody heavy chain variable region. The number of the first and last amino acid or nucleotide in each line is indicated in the left and right margins, respectively. CDRs are underlined.
- 35 Figure 2 shows the nucleotide and predicted amino acid

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sequence of mouse anti-CD38 antibody light chain variable region. The number of the first and last amino acid or nucleotide in each line is indicated in the left and right margins respectively. CDRs (underlined) were identified by comparison to known immunological sequences (Kabat et al, "Sequences of proteins of immunologic interest", US Dept of Health and Human Services, US Government Printing Office, 1987).

10 Figures 3 and 3a together show the nucleotide and predicted amino acid sequence of the humanised anti-CD38 antibody light chain cDNA. The number of the first and last amino acid or nucleotide in each line is indicated in the left and right margins, respectively. CDRs are underlined.

Figure 4 shows the nucleotide and predicted amino acid sequence of the humanised anti-CD38 antibody heavy chain cDNA. The number of the first and last amino acid or nucleotide in each line is indicated in the left and right margins, respectively. CDRs are underlined.

Figure 5 shows the configuration of the CDRHI (dark tubes) in the murine-anti-CD38 (murine residues at positions 29 and 78).

Figure 6 shows the configuration of the CDRHI (dark tubes) in the same region as Figure 5, but in a humanised construct with human residues at positions 29 and 78.

Figure 7 shows the effect of various heavy chain framework substitutions on relative binding affinity of anti-CD38 antibodies.

35 Figure 8 shows the effect of various heavy chain

framework substitutions on antibody dependent cellular cytotoxicity mediated by CD38 antibodies.

### Examples

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### Example 1

Humanisation of anti-CD38 based on a mouse antibody (AT13/5:IqGLK)

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(a) General note on methodology

Unless otherwise stated, in the medodology described below, the following standard procedures and conditions were used. Manufacturers' recommended protocols were followed where applicable.

PCR experiments (Saiki et al, Science 239:487-491, 1988) were conducted using a programmable thermal cycler (Trio Biometra). A typical 100µl reaction mix contained 2.5 units of AmpliTag polymerase (Perkin-Elmer Cetus, Beaconsfield, UK) in the buffer supplied by the manufacturer; 250µM of each of dATP, dCTP, DGTP and dTTP, amplification primers at 1 µM, and template DNA. Unless otherwise noted, the following cycle specifications were used:

step 0: 94°C for 90 seconds

step 1: 94°C for 60 seconds

30 step 2: 50°C for 60 seconds, ramping up to step 3 at a rate of 0.15°C/second

step 3: 72°C for 60 seconds, go to step 1, repeating this loop for 25 cycles

step 4: 72°C for 10 minutes.

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DNA sequencing was performed by the dideoxy method using the <u>Sequenase v2</u> system (USB, Cambridge, UK), according the manufacturer's instructions. The reaction products were separated on 8% acrylamide sequencing gels (Gel-Mix 8, BRL, Paisley, Scotland, UK).

To gel-purify DNA, one of two methods was used. fragments smaller than 175 base-pairs, the DNA was separated on a conventional high-melting point agarose gel, and the DNA recovered using the Prep-a-Gene system 10 (Bio-Rad Laboratories, Hemel Hempstead, UK). Larger fragments were purified by separation on a low-melting point agarose gel (NuSieve GTG, FMC, Rockland, ME), and the DNA recovered using Magic PCR Preps (Promega, Southampton, UK). 15

Numbering of amino-acid residues in antibody chains follows the scheme of Kabat et al ("Sequences of proteins of immunological interest", US Dept of Health and Human 😥 Services, US Government Printing Office, 1991).

- Cloning and Sequencing of AT 13/5 antibody Heavy (b) Chain...
- from a extracted was Polyadenylated RNA 25  $5 \times 10^6$  of the AT13/5 mouse hybridoma line containing using a Micro Fast Tract kit (British Biotechnology, This was converted into oligo-dT-primed - Oxford, UK). SuperScript CDNA using the single-stranded Preamplification system (BRL, Paisley, Scotland, UK). 30 Aliquots of the resulting cDNA were used in PCRs designed to separately amplify the variable region of mouse immunoglobulin heavy and light chains.
- The variable region of the heavy chain was amplified 35

according to the method of Jones & Bendig (Bio/Technology 9:88-89), using a cocktail of primers specific to the signal peptide region (MHV1-12) and one primer specific for the mouse  $\gamma$ l constant region (Mouse IgG1 heavy chain reverse primer). The resulting PCR fragment was digested with Xma I and Sal I and cloned into pUC18. Clones obtained from two independent PCR reactions were sequenced on both strands and found to be identical implying that the sequence does not contain errors introduced by the PCR process. The complete sequence of the variable region appears as Figure 1.

(c) Cloning and Sequencing of AT13/5 antibody - Light, Chain

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The sequence of the variable region of the light chain was also derived by a PCR-based cloning strategy using the same preparation of single-stranded cDNA as for the heavy chain. However, a more complex cloning and sequencing protocol was required, as the primers described by Jones & Bendig (op cit) preferentially amplify a non-productively rearranged kappa light chain from the AT13/5 cDNA. This chain arises from the fusion partner used to generate the AT13/5 hybridoma, here termed the MOPC-21 related  $V_{\rm K}$ , and is of known sequence (Carroll, WL et. al., Molecular Immunology 25:991-995; 1988).

To amplify the cDNA encoding the anti-CD38 light chain a PCR was performed using the mouse kappa light chain reverse primer described by Jones & Bendig (op cit), and a primer VK1-BACK that hybridises to the framework 1 region of most mouse kappa chains (sequences: 5'GACATTCAGCTGACCCAGTCTCCA 3'). Conditions were as described for the heavy chain amplifications above,

except that 35 cycles were used. These primers do not amplify the cDNA encoding the MOPC-21 related VK under these conditions.

An amplification fragment of the appropriate size was purified and a portion of this DNA used as the template for a second amplification (conditions as above, 30 cycles) using the light chain reverse primer and a variant of VK1-BACK containing a Hind III site (sequence: 5' GATCAAGCTTGACATTCAGCTGACCCAGTCTCCA 3'). The resulting 10 fragment was digested with Hind III and Xma I and cloned into a pUC18. Clones were sequenced on both strands by the conventional dideoxy method. Additionally, a portion of the PCR product was directly sequenced using a thermal cycling strategy (fmol system, Promgea, Southampton, UK) 15 with a primer (light chain reverse primer, as above) end- ... labelled with 32P. The sequence obtained from the cycle sequencing experiment matched exactly the sequence derived by conventional methods.

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Since this sequence was obtained from the products of two rounds of amplification, further confirmation of its accuracy was sought. The existing light chain sequence was used to design a primer that hydridises to the framework 1 region (sequence: 5' ACTAGTCGACCATCCTCTTTTCTGTTTCTCTAGGAG 3'). This was used in conjunction with the light chain reverse primer in a PCR with the following cycle definition:

30 step 0: 95°C for 120 seconds
 step 1: 95°C for 60 seconds
 step 2: 50°C for 60 seconds
 step 3: 72°C for 60 s conds, go to step 1, repeating this
 loop for 30 cycles
35 step 4: 72°C for 10 minutes

Three independent reactions were performed, and after purification, the products were digested by Xma I and Sal I, and cloned into pUC18. Several clones were sequenced by the dideoxy method. All sequences so obtained were identical to those obtained previously, confirming that the proposed light chain sequence was indeed free from PCR errors. The complete sequence of the variable region of the light chain appears as Figure 2.

10 (d) Design and construction of version 1 of the humanised antibody

Human variable domain frameworks were selected by the best-fit homology method (Lewis, AP & Crowe, JS in "Generation of Antibodies by Cell and Gene Immortalisation", Terhorst, C, Malavasi, F, Albertini, A (eds) Karger: Basel, 1993). The frameworks chosen for humanisation process were the light and heavy chain variable domains of Campath 1H (disclosed in EP-A-0328404). The humanised heavy and light chains were then constructed by a recombinant PCR technique (Lewis & Crowe, Gene 101:297-302, 1991).

### i) Light Chain

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The primers used in the humanisation process were:

- A<sub>L</sub>: 5 'GATCAAGCTTCTCTACAGTTACTGAGCACA3'
- B<sub>L</sub>: 5 'CCGATTATATATGTCCTCACTTGCCTTACAGGTGATGGTCAC3'
- 30 C<sub>L</sub>: 5'AGTGAGGACATATATAATCGGTTAACCTGGTACCAGCAGAAG3'
  - $D_L: 5$  'AGTTTCCAAACTGGTTGCACCAGAGATCAGCAGCTTTGG3'
  - EL:5'GGTGCAACCAGTTTGGAAACTGGTGTGCCAAGCAGA3'
  - F<sub>L</sub>: 5 'GTACGGATTACTCCAATACTGTTGGCAGTAGTAGGTGGC3'
  - G<sub>L</sub>: 5'CAGTATTGGAGTAATCCGTACACGTTCGGCCAAGGGACC3'
- 35 H<sub>L</sub>:5'GATCAAGCTTCTAACACTCTCCCCTGTTGA3'

Primers  $A_L$  and  $H_L$  contain Hind III sites to allow cloning of the final amplficiation product. PCRs were performed according to the following cycle specification:

5 step 0: 95°C for 120 seconds

step 1: 95°C for 60 seconds

step 2: 45°C for 60 seconds

step 3: 72°C for 60 seconds, go to step 1, repeating this

loop for 25 cycles

10 step 4: 72°C for 10 minutes

The template used in this reaction was DNA encoding the Campath 1H light chain, a construct in which the framework residues are taken from REI and the CDRs from a rat anti-human CDw52 antibody (Reichmann, L. et. al. Nature 332:323-337, 1988). The primers above are designed to wholly replace the Campath 1H sequence, leaving the AT13/5 CDRs grafted onto the REI frameworks.

- Four initial PCRs were performed using 10ng of template with the primer pairs: A<sub>L</sub> and B<sub>L</sub>, C<sub>L</sub> and D<sub>L</sub>, E<sub>L</sub> and F<sub>L</sub>, and G<sub>L</sub> and H<sub>L</sub>. The products of these reactions, AB<sub>L</sub>, CD<sub>L</sub>, EF<sub>L</sub> and GH<sub>L</sub> respectively were gel-purified and half of the amount recovered used in the second round of PCRs.

  Fragments AB<sub>L</sub> and CD<sub>L</sub> were used as template with primers A<sub>L</sub> and D<sub>L</sub> in one reaction, and fragments EF<sub>L</sub> and GH<sub>L</sub> were used as template with primers end as template with primers E<sub>L</sub> and H<sub>L</sub>. The reaction conditions were:
- 30 step 0: 95°C for 120 seconds
   step 1: 95°C for 60 seconds
   step 2: 45°C for 60 seconds
   step 3: 72°C for 90 seconds, go to step 1, repeating this
   loop for 20 cycles

The products of these reactions,  $AD_L$  and  $EH_L$ , were gelpurified and half of each DNA used as template in a final reaction with primers  $A_L$  and  $H_L$  with the reaction conditions as for the second round PCR above. The resulting product was digested with Hind III and cloned into pUC18. A clone with the predicted structure as determined by complete sequence of the insert on both strands was chosen for further manipulation. The sequence of the variable region of this construct is given as Figures 3 and 3a.

### ii) Heavy Chain

The primers used in the humanisation process were:

15 A<sub>B</sub>: 5'GATCAAGCTTTACAGTTACTCAGCACACAG3'

B<sub>R</sub>: 5'GTGGACACCATAACTGGTGAAGGTGAAGCC3'

C<sub>B</sub>: 5 'AGTTATGGTGTCCACTGGGTGAGACAGCCA3'

DH: 5 'TTGTAGTCTGTGCTTCCACCTCCACATCACTCCAATCCACTCAAG3'

E<sub>H</sub>: 5 'GAAGCACAGACTACAATGCAGCTTTCATGTCCAGAGTGACAATGCTG3'

F<sub>H</sub>: 5'GGAGTCCATCACGAAGCCGGTCGTTATCATGGATTTTGCACAATAATAGA

G<sub>H</sub>: 5 'AAATCCATGATAACGACCGGCTTCGTGATGGACTCCTGGGGTCAAGGCTC ACTAGTCACAGTCTCCTCAGCC3'

HH: 5'TAGAGTCCTGAGGGAATTCGGACAGCCGGGAAGGTG3'

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PCRs were performed according to the following cycle specification:

step 0: 95°C for 120 seconds

30 step 1: 95°C for 60 seconds

step 2: 45°C for 60 seconds

step 3: 72°C for 60 seconds, go to step 1, repeating this

loop for 25 cycles

step 4: 72°C for 10 minutes

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The template used in this reaction was DNA encoding the Campath 1H heavy chain, a construct in which the CDRs and framework residues 27 and 30 are taken from a rat antihuman CDw52 antibody (Reichmann, L et. al. op cit), and the remainder of the framework residues from NEW. primers above are designed to replace the Campath 1H CDR sequences, leaving the AT13/5 CDRs grafted onto the Campath 1H framework. Also, heavy chain residue 94 is known to be important in antigen-binding (Tempest, PR et. al., Bio/Technology, 9:260-271, 1991), so the AT13/5 10 sequence was adopted at this position. The rat sequence at residues 27 and 30 is more homologous to the AT13/5 sequence than is the unmodified NEW sequence. and  $H_{\rm H}$  contains Hind III and EcoR Additionally, primer  $G_{\text{H}}$  engineers a Spel respectively. 15 site into the framework 4 region to allow coupling to a previously prepared human C<sub>B</sub> sequence.

Four initial PCRs were performed using 10ng of template with the primer pairs:  $A_{\rm H}$  and  $B_{\rm H}$ ,  $C_{\rm H}$  and  $D_{\rm H}$ ,  $E_{\rm H}$  and  $F_{\rm H}$ , and  $G_{\rm H}$  and  $H_{\rm H}$ . The products of these reactions,  $AB_{\rm H}$ ,  $CD_{\rm H}$  were used as template with primers  $A_{\rm H}$  and  $D_{\rm H}$  in one reaction, and fragments  $EF_{\rm H}$  and  $GH_{\rm H}$  were used as template with primers  $E_{\rm H}$  and  $H_{\rm H}$ . The reaction conditions were:

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step 0: 95°C for 120 seconds

step 1: 95°C for 60 seconds

step 2: 45°C for 60 seconds

step 3: 72°C for 90 seconds, go to step 1, repeating this

. 30 loop for 20 cycles

The products of these reactions,  $AD_H$  and  $EH_H$ , were gelpurified and half of each DNA used as template in a final reaction with primers  $A_H$  and  $H_H$  with the reaction conditions as for the second round PCR above. The

resulting product was digested with Hind III and Spe I, and the fragment containing the variable region cloned into a pUC18-based vector containing the human  $C_{\rm H}$  sequence. A clone with the predicted structure as determined by complete sequencing of the insert on both strands was chosen for further manipulation.

(e) Eukaryotic expression of version 1 of the humanised antibody

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Humanised AT13/5 heavy and light chains were cloned into eukaryotic expression vectors under human β actin promoters. The heavy and light chain plasmids were transiently expressed in B11 CHO cells by cotransfection of the two plasmids using <u>Transfectam</u> (Promega, Southampton, UK). Culture supernatants were assayed for human IgG by ELISA, and tested for CD38-binding activity by FACS analysis using the CD38-positive B-cell line Wien 133.

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Although the culture supernatants/contained significant amounts of human IgG, no anti-CD38 activity could be detected by FACS, even when supernatants were concentrated 10-fold. This result suggests that simple grafting of the CDRs from AT13/5 onto the Campath 1H and REI human frameworks is insufficient to transfer the antibody specificity. A series of framework changes were therefore undertaken in order to restore CD38-binding activity.

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### (f) Framework changes

Since most of the framework residues previously shown to be important in r storing antigen binding are in th heavy chain variable region, it was decided to focus on

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this part of the antibody. Additional cotransfection of the humanised light chain with a chimaeric heavy chain construct (mouse heavy variable region fused to human  $C_{\rm H}$ ), produced active antibody (hereafter termed hybrid antibody) that bound CD38 with an affinity comparable to that of the original mouse antibody. The region with the lowest homology between the human frameworks used and the original mouse sequence is also close to some residues of known importance. This region, just downstream of the CDR3 sequence was chosen for mutagenesis.

Heavy chain residues 67 to 71 inclusive and 73 were grafted from the mouse antibody onto the humanised heavy chain using recombinant PCR. The primers used were as follows:

An: sequence as above

In: 5'GTTGTCCTTGGTGATGTTCAGTCTGGACATGAAAGCTGC3'

J<sub>B</sub>: 5'CTGAACATCACCAAGGACAACAGCAAGAACCAGTTCAGC3'

20  $H_B$ : sequence as above.

Two initial PCRs were performed using 10ng of version 1 humanised heavy chain template with the primer pairs:  $A_{\rm H}$  and  $I_{\rm H}$  and  $J_{\rm H}$  and  $H_{\rm H}$ . The products of these reactions,  $AI_{\rm H}$  and  $JH_{\rm H}$  respectively, were gel-purified and half of the recovered DNA used in a second round of PCR with primers  $A_{\rm H}$  and  $H_{\rm H}$  to generate version 2 of the humanised heavy chain variable region. This was cloned, sequenced, transferred to the expression system, and then transiently co-expressed with the humanised light chain construct as above. Once again, culture supernatant from transfected CHO c 11s produced human IgG as determined by ELISA, but no CD38-binding activity could be detected by FACS analysis.

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A further round of mutations based on both version 1 and version 2 of the humanised heavy chain were then produced by a method identical to that described above. A total of six version 3 heavy chains were produced in which the following heavy chain framework residues were grafted from the mouse sequence onto one or other humanised sequence:

	Antibody	Y Template for	Grafted residues	Primers used
10	٠	mutagenesis		
	h3J	version 1	28,29	K <sub>H</sub> , L <sub>H</sub>
	h3K	version 2	28,29	$K_{\rm H}$ , $L_{\rm H}$
	h3L	version 1	76	M <sub>H</sub> , O <sub>H</sub>
	h3M	version 2	76	$N_{\rm H}$ , $O_{\rm H}$
15]	h3N	version 1	28,29,76	$K_H, L_H, M_H, O_H$
	h30	version 2	28,29,76	$K_H, L_H, N_R, O_B$

Additionally, all constructions used primers  $A_{\text{H}}$  and  $H_{\text{H}}$ . The primer sequences used were:

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A<sub>H</sub>: sequence as above

H<sub>H</sub>: sequence as above

KH: 5'ACTGGTTAACGAAAAGCCAGACACGGTGCAGGTCAG3'

Lm: 5'GGCITTTCGTTAACCAGTTATGGTGTCCACTGGGTG3'

5 M<sub>B</sub>: 5'AAATTGCCGTTTCGAAGTGTCTACCAGCATTGTCAC3'

N<sub>B</sub>: 5'AAATTGCCGTTTCGAATTGTCCTTGGTGATGTTCAG3'

OH: 5'TTCGAAACGGCAATTTAGCTTGAGACTCAGCAGC3'

Heavy chain constructs containing the expected sequence were transferred into mammalian expression vectors, and cotransfected with the humanised light chain construct into CHO cells, as above. Tissue culture supernatants containing human IgG as determined by ELISA were assayed for CD38-binding activity by FACS. Constructs h3K and h3O showed antigen-binding in this assay though with less

activity than the hybrid antibody (see Fig. 7).

Method for changing framework residues at positions 29 and 78

In order to establish why h30 showed less activity than the hybrid antibody further sequences analysis suggested potential problems with positions 29 and 78 in the heavy chain.

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Having identified mutations to be made in the heavy chain framework regions, these can be produced by a variety of examples being site-directed methods: standard mutagenesis, recombinant PCR and gene synthesis using oligonucleotides. In the case of the anti-CD38 heavy chain VH, recombinant PCR was used to introduce murine residues at positions 28-29 and 78 sequentially.

A human anti-CD38 heavy chain VH already incorporating m murine residues at positions 27, 30, 67, 68, 69, 70, 71, 20

73 and 94 (Version 2 as described in (f) above) was used as template for the first round of mutagenesis. This was amplified with the following PCR primers in two separate

reactions:

-Primer A: 5'GATCAAGCTTTACAGTTACTCAGCACAG3' 25

Primer B: 5'ACTGGTTAACGAAAAGCCAGACACGGTGCAGGTCAG3'

Primer C: 5'GGCTTTTCGTTAACCAGTTATGGTGTCCACTGGGTG3'

Primer D: 5'TAGAGTCCTGAGGGAATTCGGACAGCCGGGAAGGTG3'

In primers B and C, the triplets encoding the murine residues at positions 28 and 29 are underlined. 30 first reaction, the template was amplified with primers In the second reaction, the template was amplfified with primers C and D. The products of the two reactions were purified, mixed, and amplified with primers A and D. The reaction product was purified,

cleaved with Hind III and SpeI, and the 450 base-pair fragment encoding the VH cloned into a variant of pUC18 containing a human vl cDNA cassette (Sime et al, 1993; J. Immunol, 151:2296). Clones were sequenced to ensure correct introduction of the murine residues at positions 28 and 29.

A clone incorporating these changes was then used as template for a second round of recombinant PCR 10 mutagenesis to introduce the murine residue at position 78. A procedure identical to that described above was followed, except that primers B and C were replaced by primers E and F respectively, which contain a triplet (underlined) that incorporates the murine residue at position 78.

Primer E: 5'AACCAGGTGAGCTTAAGACTCAGCAGCGTGACA3'
Primer F: 5'TCTTAAGCTCACCTGGTTCTTGCTGTTGTCCTT3'

- The resulting heavy chain (see Fig. 4) when co-expressed with the humanised light chain (see Fig. 3) produces humanised anti-CD38,h3S.
- (h) Eukaryotic expression of functional humanised
  25 antibody

To creat clonal cell lines for further characterisation, plasmids encoding the humanised h3S heavy chain and the chimaeric heavy chain were separately co-transfected with the humanised light chain into B11 CHO cells.

### Example 2

#### Biological activity

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# (a) CD38 Binding Studies

(i) Effect of various heavy chain framework substitutions on relative binding affinity of anti-CD38 antibodies.

Binding was assessed by FACS staining of CD38 positive cells.

Heavy chains incorporating one or more of mouse framework residues were created as described above and combined with the humanised light chain to make antibodies which were assayed for binding to CD38, with the following results.

15	Construct	66-73	28/29	78	Binding
	h1	_	-	-	· • –
	h2	+	<u> </u>	- "	-
•	h3J · ·	_	+	-	·
•	h3K	. +	+	. • •	· + ,
20	h3S	*** • • • • • • • • • • • • • • • • • •	* +	+ ,	++

In this table, + denotes that the murine framework residue is incorporated into the humanised antibody at the indicated position, - denotes that the human residue remains.

#### Discussion

According to computer modelling studies the change of the 66-73 region back to mouse framework causes the humanised CDRH2 to adopt a similar conformation to that of the mouse antibody. However, as the construct h2 shows, this is insuffici nt to obtain binding. The model also suggests that in the mouse anti-CD38 antibody, positions 29 and 78 are occupied by small residues, whose side-

chains pack neatly together allowing CDRH1 to adopt the correct configuration for antigen binding. In the humanised constructs h1 and h2, the side chains are unable to pack together in this fashion, being much larger, and so distort CDRH1, preventing antigen binding. This aspect of the model is illustrated in Figures 5 and 6 (attached). Figure 5 shows the configuration of CDRH1 (dark tubes) in the murine anti-CD38. In Figure 6 showing the same region in a humanised construct with human residues at positions 29 and 78, the extra bulk of these side chains has clearly resulted in a distortion of the CDRH1 conformation.

Partial relief of this effect can be obtained by using the murine residue at position 29 and the human residue at position 78, though the resulting antibody shows markedly reduced function. Use of murine residues at both positions 29 and 78 restores activity, as evidenced by the data for the h3S construct.

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(ii) Anti-CD38 heavy chain variable regions were fused to human γl constant region and coexpressed in CHO cells with humanised anti-CD38 light chain. CD38-binding activity is expressed normalised to the signal obtained using a saturating dose of hybrid antibody (mouse VH) in the same experiment.

Results are shown in Figure 7 where:

- Humanised antibody with murine residues at 28,29 and 78
  - ▲ Humanised antibody with murine residues at 28,29 and 76

- Humanised antibody with murine residues at 28,29
- Hybrid antibody
- In addition to the above substitutions, all humanised heavy chains contained murine framework residues at positions 27, 30, 67, 68, 69, 70, 71, 73 and 94. These alone are insufficient to obtain detectable binding by FACS.

These results demonstrate the critical importance of the small residues at positions 29 and/or 78 in obtaining full humanised heavy chain activity. They also demonstrate the specific nature of the interaction, in that a murine residue at position 76 close to position 78 was unable to restor activity.

(b) Effect of various heavy chain framework substitutions on antibody-dependent cellular cytotoxicity mediated by CD38 antibodies.

Antibody-dependent cellular cytotoxicity is normally assessed by one of several label-release techniques, In one such assay, 104 well-known in the literature. target cells (Wien 133) were labelled with europium and 25 then exposed to freshly prepared human peripheral blood antibody in the presence of effector:target ratio of 50:1. Lysis was estimated by detecting release of europium after 4 quantitated by reference to control reactions without 30 lymphocytes antibody or peripheral blood detergent such as Triton-X100.

The effect of framework substitutions on the lytic pot ntial of humanised anti-CD38 monoclonals was examined

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in label-release assay. Wien 133 target cells were loaded with label (either 51Cr or Eu) and then exposed to freshly prepared human peripheral blood mononuclear cells in the presence of varying amounts of anti-CD38 antibody. Cytotoxicity is expressed as the proportion of total releasable label liberated by antibody treatment.

Results are shown in Figure 10 where:

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- A Humanised antibody with murine residues at 28,29 and 78
- Humanised antibody with murine residues at 28,29 and 76
- 15 Hybrid antibody

These results show that the combination of framework changes at positions 29 and 78 confer full activity on the humanised heavy chain for cytotoxic function.

Although incorporation of a small murine residue at position 29 results in some binding activity (Figure 7), this is insufficient to achieve full effector function.

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### SEQUENCE LISTING

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- (ii) TITLE OF INVENTION: ANTIBODIES
- (iii) NUMBER OF SEQUENCES: 46
  - (iv) COMPUTER READABLE FORM:
    - (A) MEDIUM TYPE: Floppy disk

(B) COMPUTER: IBM PC compatible

(C) OPERATING SYSTEM: PC-DOS/MS-DOS

(D) SOFTWARE: PatentIn Release #1.0, Version #1.30 (EPO)

# (2) INFORMATION FOR SEQ ID NO: 1:

### (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 454 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

#### (ix) FEATURE:

(A) NAME/KEY: CDS

(B) LOCATION:1..453

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 1:

GGT CGA CTG GCT GTG TTA GCG CTG CTC TTC TGC CTG GTG ACA TTC CCA
Gly Arg Leu Ala Val Leu Ala Leu Leu Phe Cys Leu Val Thr Phe Pro

1 5 10 15

AGC TGT GTC CTG TCC CAG GTG CAG CTG AAG CAG TCA GGA CCT GGC CTA 96
Ser Cys Val Leu Ser Gln Val Gln Leu Lys Gln Ser Gly Pro Gly Leu
20 25 30

GTG CAC CCC TCA CAG AGC CTG TCC ATA ACC TGC ACA GTC TCT GGT TTC

Val His Pro Ser Gln Ser Leu Ser Ile Thr Cys Thr Val Ser Gly Phe

35

40

45

TCA TTA ACT AGT TAT GGT GTC CAC TGG GTT CGC CAG TCT CCA GGA AAG 19

											•						
Ser		Thn	Ser	Tyr	Gly	Va 1 55	His	Trp	Va 1	Arg	G1n 60	Ser	Pro	Gly	Lys		
	50					33											•
GGT	CTG	GAG	TGG	CTG	GGA	GTG	ATG	TGG	-AGA	GGT	GGA	AGC	ACA	GAC	TAC		240
Gly	Lau	Glu	Tro	Leu	G1v	Val	Het	Trp	Arg	G1y	Gly	Ser	Thr	Asp	Tyr		
65		• • •		-	70			·		75	•				80	•	•
Ų,						÷		•									
TAA	GCA	GCT	TTC	ATG	TCC	AGA	CTG	AAC	ATC	ACC	AAG	GAC	AAC	TCC	AAG	•	288
Asn	Ala	Ala	Phe	Met	Ser	Arg	Leu	Asn	Ile	Thr	Lys	Asp	Asn	Ser	Lys	•	
				85		٠			90					95	i .: .		
		•				8											
CGC	CAG	GTT	ттс	ш	AAA	ATG	AAC	AGT	CTA	CAA	GCT	GAT	GAC	ACT	GCC	•	336
Ara	Gln	Va 1	Phe	Phe	Lys	Het	Asn	Ser	Leu	G1n	Ala	Asp	Asp	Thr	· Ala		
			100	-	,			105					110				•
							,										*.
ATA	TAC	TAC	TG1	GCC	: AAA	TCG	ATG	ATT	r AC	ACG	GGC	मा	GTT	ATE	GAC		384
Ile	Tyr	Tyı	r Cys	s Ala	Lys	Ser	Met	: 11	e Thi	r Thi	r Gly	Phe	e Va	1 He	t Asp		
		11!					120					129	5	٠			
			•					1,		•							
TCC	TGO	G GG	T CA	A GG	A ACC	TC/	A GTO	: AC	C GT	C TC	C TC	A GCC	CAA	A ACI	G ACA	*	432
Ser	Tri	o G1	v Gl	n G1	y Thi	r Sei	r Va	1 Th	r Va	1 Se	r Se	r Al	a Ly	s Th	r Thr		:
•	130		, ,		•	13					14		•				
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CCC	. cc	A TC	T GT	C TA	T CC	A CT	GG	r		1 .1							454
				1 Ty													
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		-				-					•						

# (2) INFORMATION FOR SEQ ID NO: 2:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 151 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 2: Gly Arg Leu Ala Val Leu Ala Leu Leu Phe Cys Leu Val Thr Phe Pro <sub>t.</sub> 10 Ser Cys Val Leu Ser Gln Val Gln Leu Lys Gln Ser Gly Pro Gly Leu 20 25 Val His Pro Ser Gln Ser Leu Ser Ile Thr Cys Thr Val Ser Gly Phe 35 Ser Leu The Ser Tyr Gly Val His Trp Val Arg Gln Ser Pro Gly Lys Gly Leu Glu Trp Leu Gly Val Met Trp Arg Gly Gly Ser Thr Asp Tyr 65 -70 75 Asn Ala Ala Phe Met Ser Arg Leu Asn Ile Thr Lys Asp Asn Ser Lys 85 95 Arg Gln Val Phe Phe Lys Het Asn Ser Leu Gln Ala Asp Asp Thr Ala 100 105 Ile Tyr Tyr Cys Ala Lys Ser Net Ile Thr Thr Gly Phe Val Met Asp 115 120 125 Ser Trp Gly Gln Gly Thr Ser Val Thr Val Ser Ser Ala Lys Thr Thr 130 135 140

(2) INFORMATION FOR SEQ ID NO: 3:

150

Pro Pro Ser Val Tyr Pro Leu

# (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 454 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

# (iv) ANTI-SENSE: YES

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 3:

CCAGCTGACC GACACAATCG CGACGAGAAG ACGGACCACT GTAAGGGTTC GACACAGGAC	60
AGGGTCCACG TCGACTTCGT CAGTCCTGGA CCGGATCACG TGGGGAGTGT CTCGGACAGG	120
TATTGGACGT GTCAGAGACC AAAGAGTAAT TGATCAATAC CACAGGTGAC CCAAGCGGTC	180
AGAGGTECTT TECCAGACET CACEGACECT CACTACACET ETECACETTE GTGTETGATG	240
TTACGTCGAA AGTACAGGTC TGACTTGTAG TGGTTCCTGT TGAGGTTCGC GGTCCAAAAG	300
AAATTITACT TGTCAGATGT TCGACTACTG TGACGGTATA TGATGACACG GTTTAGCTAC	360
TAATGCTGCC CGAAACAATA CCTGAGGACC CCAGTTCCTT GGAGTCAGTG GCAGAGGAGT	420
CGGTTTTGCT GTGGGGGTAG ACAGATAGGT GACC	454

# (2) INFORMATION FOR SEQ ID NO: 4:

# (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 364 base pairs

(B) TYPE: nucleic acid

. 44

(C) STRANDEDNESS: double (D) TOPOLOGY: linear

# (ix) FEATURE:

235

(A) NAME/KEY: CDS
(B) LOCATION:1..363

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 4:

GAC	ATT	CAG	CTG	ACC	CAG	TCT	CCA	TCC	TCC	ш	TCT	GTT	TCT	CTA	GGA	÷	48
Asp	Ile	Gln	Leu	Thr	Gln	Ser	Pro	Ser	Ser	Phe	Ser	Val	Ser	Leu	Gly		
•			155			e		160			•		165	,	- '	•	
					•		•	••	- :				•				
GAC	AGA	GTC	ACC	ATT	ACT	TGC	AAG	GCA	AGT,	GAG	GAC	ATA	TAT	AAT	CGG	*	96
Asp	Arg	Val	Thr	Ile	Thr	Cys	Lys	Ala	Ser	Glu	Asp	Ile	Tyr	Asn	Arg		
		170					175		0			180					
					•	,		- 000			•		•		•	• • •	
TTA	ACC	TGG	TAT	CAG	CAG	AAA	CCA	GGA	AAT	GCT	ССТ	AGG	CTC	TTA	ATA		144
															Ile	* • • • • • • • • • • • • • • • • • • •	
٠	185	-				190					195					• • • • •	
		• •	• , , • •	. •				<i></i>	•	٠.				· .			
TCT	GGT	GCA	ACC	AGT	TTG	GAA	ACT	GGG	GTT	CCT	TCA	AGA	TTC	AGT	GGC		192
	•								-						Gly		
200					205					210		3			215		
									٠.							*	
AGT	GGA	тст	GGA	AAG	GAT	TAC	ACT	CTC	AGC	ATT	ACC	AGT	CTT	CAG	ACT		240
								. ;	, ,						Thr	4	
		, 0,00	3	220	_				225			-		230		-	*
	*	- 50		LLU					223					230	٠.		
GAA	CAT	СТТ	CCT	ACC	TAT	TAC	TOT	CAA	CAC	Tay	TCC	ACT			TAC		200
						*			-			,			TAC		288
Giu	ASP	val	Ala	Inr	i yr	lyr	Lys	GIN	GIN	ıyr	ırp	) Ser	ASI	Pro	Tyr		

240

364

45

ACG TTC GGA GGG GGG ACC AAG CTG GAA ATA AGA CGG GCT GAT GCT GCA Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Arg Arg Ala Asp Ala Ala 260 255 250 CCA ACT GTA TCC ATC TTC CCA CCA TCC A Pro Thr Val Ser Ile Phe Pro Pro Ser 265 270 (2) INFORMATION FOR SEQ ID NO: 5: (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 121 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear (ii) MOLECULE TYPE: protein (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 5: Asp Ile Gln Leu Thr Gln Ser Pro Ser Ser Phe Ser Val Ser Leu Gly 15 10 1 Asp-Arg-Val Thr Ile Thr Cys Lys Ala Ser Glu Asp Ile Tyr Asn Arg 20 Leu Thr Trp Tyr Gln Gln Lys Pro Gly Asn Ala Pro Arg Leu Leu Ile 40 35 Ser Gly Ala Thr Ser Leu Glu Thr Gly Val Pro Ser Arg Phe Ser Gly 60 55 50 Ser Gly Ser Gly Lys Asp Tyr Thr Leu Ser Ile Thr Ser Leu Gln Thr 80 75 -70 65

Glu Asp Val Ala Thr Tyr Tyr Cys Gln Gln Tyr Trp Ser Asn Pr Tyr

85

90

95

Thr Phe Gly Gly Gly Thr Lys Leu Glu Ile Arg Arg Ala Asp Ala Ala 100 105 110

Pro Thr Val Ser Ile Phe Pro Pro Ser 115 120

#### (2) INFORMATION FOR SEQ ID NO: 6:

#### (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 364 base pairs.

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

#### (iv) ANTI-SENSE: YES

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 6:

CTGTAAGTCG ACTGGGTCAG AGGTAGGAGG AAAAGACAAA GAGATCCTCT GTCTCAGTGG 60

TAATGAACGT TCCGTTCACT CCTGTATATA TTAGCCAATT GGACCATAGT CGTCTTTGGT 120

CCTTTACGAG GATCCGAGAA TTATAGACCA CGTTGGTCAA ACCTTTGACC CCAAGGAAGT 180

TCTAAGTCAC CGTCACCTAG ACCTTTCCTA ATGTGAGAGT CGTAATGGTC AGAAGTCTGA 240

CTTCTACAAC GATGGATAAT GACAGTTGTC ATAACCTCAT TAGGCATGTG CAAGCCTCCC 300

CCCTGGTTCG ACCTTTATTC TGCCCGACTA CGACGTGGTT GACATAGGTA GAAGGGTGGT 360

47

AGGT				364
(2) INFORMATION FOR SEQ ID NO:	7:		· ·	
(i) SEQUENCE CHARACTERISTI	CS:			
(A) LENGTH: 746 base				+
(B) TYPE: nucleic act			· E = *	
(C) STRANDEDNESS: dou	4			
(D) TOPOLOGY: linear				
				. *
	* *	*		
(ix) FEATURE:				† *
(A) NAME/KEY: CDS		* . * .		
(B) LOCATION:3737		*		
			** *	
(xi) SEQUENCE DESCRIPTION:	SEQ IÓ NO: 7	<b>:</b>	*	The state of the s
AA GCT TCT CTA CAG TTA CTG AGG	ACA CAG GAC	CTC ACC ATG GGA	TGG	47
Ala Ser Leu Gln Leu Leu Se	r Thr Gln Asp	Leu Thr Met Gly	y Trp	
125	130	13	5	•
AGC TGT ATC ATC CTC TTC TTG G	TA GCA ACA GC	T ACA GGT GTC CA	AC TCC	95
Ser Cys Ile Ile Leu Phe Leu V		a Thr Gly Val H	is Ser	المستأثية
140	145	₹ <b>150</b>		
GAC ATC CAG ATG ACC CAG AGC C	CA ACC ACC CT	G AGC GCC AGC G	TG GGT	143
Asp Ile Gln Met Thr Gln Ser F	orn Ser Ser Le	u Ser Ala Ser V	al Gly	
	160 Sei Sei E	165		
122	•			

GAC AGA GTG ACC ATC ACC TGT AAG GCA AGT GAG GAC ATA TAT AAT CGG

Asp Arg Val Thr Ile Thr Cys Lys Ala Ser Glu Asp Ile Tyr Asn Arg

175

										144						
TTA	ACC	TGG	TAC	CAĢ	CAG	AAG	CCA	GGT	AAG	GCT	CCA	AAG	CTG	CTG	ATC	239
Leu	Thr	Trp	Tyr	Gln	G1n	Lys	Pro	Gly	Lys	Αla	Pro	Lys	Leu	Leu	Ile	
185					190				,	195	-			٠.	200	
TCT	GGT	GCA	ACC	AGT	TTG	GAA	ACT	GGT	GTG	CCA	AGC	AGA	TTC	AGC	GGT	287
															Gly	
	3			205	-				210					215		• • • • • • • • • • • • • • • • • • • •
								÷ .								
400	CCT	ACC	CCT	400	CAC	TTC	400	TTC	. ACC	ATC	ACC	ACC	CTC	CAG	CCA	225
			GGT								•	, 5			7	335
5er	Gly	Ser			ASP	rne	ınr		ınr	116	Ser	Ser			Pro	- :
			220					225		•			230	,		
		-								- `				. :	· •	
GAG	GAC	ATC	GCC	ACC	TAC	TAC	TGC	CAA	CAG	TAT	TGG	AGT	AAT	CCG	TAC	383
G1u	Asp	Ile	Ala	Thr	Tyr	Tyr	Cys	Gln	Gln	Tyr	Trp	Ser	Asn	Pro	Tyr	
		235	; ,				240				•	245	ī			
																*
ACG	TTO	GGC	CAA	GGG	ACC	AAG	GTG	GAA	ATC	AAA	CGA	ACT	GTG	GCT	GCA	431
Thr	• Phe	e G1v	Gln	Gly	Thr	Lys	Val	Glu	Ile	Lys	Arc	Thr	Va1	Ala	Ala	•
	250	, , , , •				255		100			260				•	
								•	: 5.							
CC	· A TCI	GTO	: TTC	ATO	TTC	CCG	CCA	тст	GAT	GAG	CAG	TTG	AAA	TCT	GGA	479
	2					•						* 1	٠.		Gly	
26		•			270				, nop	275			,.		280	
20:					2/0	- 30			•	2/-	•	- 8-			200	•
	- 00	: 														E97
															GCC	527
Th	r Ala	a Sei	r Val			Leu	Leu	ı Asn			e lyi	r Pro	) Ar		u-Ala -	•
•				28	5 · ·	•			290	)				29	•	
	•			٠.							: :		4			2
AA	A GT	A CA	G TG(	G AAI	G .GTO	GAT	AAC	GCC	CTC	CAA	A TC	GGT	, AAI	CTC	CAG	575
Ly	s Va	1 - G1:	n Trị	p Ly	s Va	l Ası	Asi	i Ala	Le	ı Glı	n Se	r G1;	y .As	n Se	r Gln	- I
			300	0			2	305	5		•	· ·	31	0	v *v	•
·.								,		-						
GA	G AG	T GT	C :AC	A. GA	G CA	G GA	AGO	CAAC	GA	CAG	CAC	C TA	CAG	C CT	C AGC	62:
		*	-						•						u Ser	
		21		- • •	_ +1	,	321	-				32			•	•

719

											49							
AGC Ser	٠ ٦	\CC  hr  330	L	rG eŭ	ACG Thr	CTG Leu	AGC Ser	AAA Lys 335	Ala	GAC Asp	TAC Tyr	GAG Glu	AAA Lys 340	CAC	Lys	GTC Val	TAC Tyr	
GC( A1:	a (	TGC Cys	G. - G	AA Tu	GTC Val	ACC	CAT His		GGC Gly	CTG Leu	AGC Ser	TCG Ser 355	Pro	GTC Val	ACA Thr	AAG Lys	AGC Ser 360	
							u Cy:	TAG	AAGC	<b>:TT</b>	*****	. · · · · · · · · · · · · · · · · · · ·	2		,	**		
(2	2)	IN						Q ID ARAC			S:	0	0.50					
-		•		(	(A) (B)	LENG Type	TH:	245 iino ': li	amin acid	o ac				•	,			. (3)
								: pr			•	NO:					: .	· ,
A	_	. Se	er	Le	eu G	ln L	eu L	eu S	er Ti	hr G		sp Le	eu Tł	ır Me	et Gi	ly Tr	p Sei	 r 
(	Çy:	s I	1e		le L	-	he Ļ	eu V	al A		25		•		al H	30	er As	P
:	11	e G	iln		et 1 35	hr (	iln S	er P						la S			ly As	P
	Ar	g \	/a 1 50		hr :	(le	Thr (	ys l	.ys <i>l</i> 55	Ala S	er 0	ilu. A	sp I	le T 60	yr A	sn A	rg Le	<b>≥u</b>

Thr 65	Trp	Tyr	Gln	Gln	Lys 70	Pro	Gly	Lys	Ala	Pro 75	Lys	Leu	Leu	Ile	Ser 80
G1y	Ala	Thr	Ser	Leu 85	GTu	Thr	Gly	Va 1	Pro 90		Arg	Phe	Ser	G1y 95	Ser
G1v	Ser	61v	Thr	Asn	Phe	The	Pho	Thr	Ila	San	San	Lou	Cla		C1
<b></b>			100	,	7110		r,iic	105		JE1	JEI	Leu	110		Giu
Asp	Ile		Thr	Tyr	Tyr	Cys		G1n	Tyr	Trp	Ser		Pro	Tyr	Thr
*	• •	115					120	٠		2 °		125			
Phe	Gly 130	Gln	Gly	Thr	Lys	Va 1 135	Glu	Ile	Lys		Thr 140	Val	Ala	Ala	Pro
Ser 145	Va 1	Phe	Ile	Phe	Pro 150		Ser	Asp		Gln 155		Lys	Ser	Gly	Thr 160
•	Ser	Val	V.1	Cve		•.	<b>A</b> = <b>T</b>	<b>A</b> = <b>-</b>					61		
714	Jei	<b>V</b> Q.1	Vai	165		Leu	VƏII	VOII	170		Pro	Arg	GIU	175	Lys
Va 1	G1n	Trp	Lys 180		Asp	Asn	Ala	Leu 185		Ser	Gly	Asn	Ser 190	1	Glu
Ser	Val	Thr 195		Gln	Asp	Ser	Lys 200		Ser	Thr	Tyr	Ser 205		Ser	Ser
Thr	L <b>e</b> u 210	Thr		Ser	-	Ala 215	•	Tyr	Glu	Lys	His 220		Val	Tyr	Ala
Cys 225	G Tu	Val	Thr	His	G1n 230	-	Leu	Ser	Ser	Pro 235	*	Thr	Lys	Ser	Phe 240
A = ==		. 61.	63	. • •	,		•		٠,				. :		٠.

# (2) INFORMATION FOR SEQ ID NO: 9:

# (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 746 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: double

(D) TOPOLOGY: linear

## (iv) ANTI-SENSE: YES

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 9:

	,		* :			
TTCGAAGAGA	TGTCAATGAC	TCGTGTGTCC	TGGAGTGGTA	CCCTACCTCG	ACATAGTAGG	60
AGAAGAACCA	TCGTTGTCGA	TGTCCACAGG	TGAGGCTGTA	GGTCTACTGG	GTCTCGGGTT	120
CGTCGGACTC	GCGGTCGCAC	CCACTGTCTC	ACTGGTAGTG	GACATTCCGT	TCACTCCTGT	180
ATATATTAGC	CAATTGGACC	ATGGTCGTCT	TEGGTCEATT	CCGAGGTTTC	GACGACTAGA	240
GACCACGTTG	GTCAAACCTT	TGACCACACG	GTTCGTCTAA	GTCGCCATCG	CCATCGCCAT	300
GGCTGAAGTG	GAAGTGGTAG	TCGTCGGÁGG	TCGGTCTCCT	GTAGCGGTGG	ATGATGACGG	. 360
TTGTCATAAC	CTCATTAGGC	ATGTGCAAGC	CGGTTCCCTG	GTTCCACCTT	TAGTTTGCTT	420
GACACCGACC	TGGTAGACAG	AAGTAGAAGG	GCGGTAGACT	ACTCGTCAAC	TTTAGACCTT	480
GACGGAGACA	ACACACGGAC	GACTTATTGA	AGATAGGGTO	: TCTCCGGTTT	CATGTCACCT	540
TCCACCTAT	r GCGGGAGGTT	AGCCCATTG	GGGTCCTCTC	ACAGTGTCT	GTCCTGTCGT	600

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AGTTGTCCCC TCTCACAATC TT	CGAA	*	ć,.	*			746
(2) INFORMATION FOR SEQ	ID NO: 10:		*				
(i) SEQUENCE CHARAC	TERISTICS:	1			9		
(A) LENGTH: 43		·			•		
(B) TYPE: nuc	₹						
(C) STRANDEDNI							
(D) TOPOLOGY:						: '	
				- , , ·			
· · ·					**		
				•			
(ix) FEATURE:		* . ***			· ja		•
(A) NAME/KEY:	CDS		•				
(B) LOCATION:	314	•					
		+					
(ix) FEATURE:			. 4	*		*	
(A) NAME/KEY:	CDS			3 7		•	
(B) LOCATION:	18434	•				e e e e e e e e e e e e e e e e e e e	
4	*	•					-
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(xi) SEQUENCE DESCR	IPTION: SE	Q ID NO:	10:				
+	*	•				0. 2.	
AA GCT TTA CAG TTA CNC	AGC ACA CA	G GAC CTC	ACC ATG	GGA TG	G AGC		- 4
Ala Leu Gln Leu	Ser Thr G1	n Asp Leu	Thr Met	Gly Tr	p Ser		8
	1	5			10	,	
•						_	

				1,141					•									
GTC	CAA	C1	rG	CAG	GAG	AGC	GGT	CCA	GGT	CTT	GTG	AGA .	CCT	AGC .	CAG	ACC	,	143
Va 1	Gln	Le	eu	Gln	Glu	Ser	Gly	Pro	Gly	Leu	Val	Arg	Pro	Ser	Gln	Thr		
		-	,	30				·	35					40	:			
• (					•			•				- Y -						•
CTG	AGC	: C	TG	ACC	TGC	ACC	GTG	TCT	GGC	TTT	TCG	TTA	ACC	AGT	TAT	GGT,		191
Leu	Sei	· L	eu	Thr	Cys	Thr	Val	Ser	Gly	Phe	Ser	Leu	Thr	Ser	Tyr	Gly		
			45					50	•				55				· ·	
•				,		f .	•						,	. 8	-			•
GTO	CA	C T	GG	GTG	AGA	CAG	CCA	CCT	GGA	CGA	GGT	CTT	GAG	TGG	ATT	GGA		239
Val	Hi	s T	rp	Va1	Arg	, GÌn	Pro	Pro	Gly	Arg	Gly	Leu	Glu	Trp	Ile	Gly	: "	
		0					65					70		. ,		:'		
					•										•			
GT	G AT	G 7	rgg	AGA	GGT	GGA	AGC	ACA	GAC	CTAC	TAA:	GCA	GCT	TTC	: ATE	TCC		287
٧a	1 Me	t i	Trp	Arg	G1:	y Gly	/ Ser	Thi	r As	p Ty	r: Asr	sfå r	s Ala	Pho	e He	t Ser		
7						80		* *			8	-			-	90	-	
							3									1.0		
AG	A CT	G.	AAC	: AT	CAC	C AÀ	G GAC	AA.	C AG	CAA	G AA	CAG	GTO	G AG	CIT	A AGA	•	335
Ar	g Lo	eu	Ası	ı Il	e Th	r Ly	s As	p Ās	n Se	r Ly	s As	n Gli	n Va	1 Se	r Le	u Arg		
					_	5		:		10					10	5		***
				. 7			27						• •					202
CT	C A	GC	AG	C GT	G AC	A GC	C GC	C GA	C AC	C GC	G GT	C TA	T TA	TTG	T GC	A AAA	-	383
Le	eu S	er	Şe	r Va	1 Th	ır Al	a Al	a As	p Tł	nr Al	a Va	1 Ty	т Ту	r Cy	S AI	a Lys		
				11	.0:_		*	<b>-</b>	1.	15	<u>.</u>			12	20 .			/ <del>-</del> -
-			-	: .		0	-									C TCA	- :	431
T	CC A	TG	AT	A AC	G A	CC GC	C TT	C GT	rg A	TG G/	AC TO	C TG	iG GG	T CA	IA GI	C TCA		
S	er P	let,	11	e Ti	ir T	hr 'G'	ly Pi			et A	sp Se	er II			in G	ly Ser		
		1	12	25				1	30			· .	1.	35			•	
		•	•							•			•					436
_	TA	T:	•		٠,									٠,	98.		·	, 700

(2) INFORMATION FOR SEQ ID NO: 11:

Leu

(i) SEQUENCE CHARACTERISTICS:
(A) LENGTH: 4 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 11:

Ala Leu Gln Leu

## (2) INFORMATION FOR SEQ ID NO: 12:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 139 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 12:

Ser Thr Gln Asp Leu Thr Met Gly Trp Ser Cys Ile Ile Leu Phe Leu

1 5 10 15

Val Ala Thr Ala Thr Gly Val His Ser Gln Val Gln Leu Gln Glu Ser
20 25 30

Gly Pro Gly Leu Val Arg Pro Ser Gln Thr Leu Ser Leu Thr Cys Thr 35 40 45

Val Sér Gly Phe Ser Leu Thr Ser Tyr Gly Val His Trp Val Arg Gln
50 55 60

Pr Pro Gly Arg Gly Leu Glu Trp Ile Gly Val Met Trp Arg Gly Gly

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65					70	-				75		• ' .		• • .	80			
Ser	Thr	Asp	Tyr	Asn 85	Ala	Ala	Phe	Met	Ser 90	Arg	Leu	Asn	Ile	Thr 95	Lys		- (4)	
Asp	Asn	Ser	Lys 100		Gln	Va 1	Ser	Leu 105		Leu	Ser	Ser	Val 110	Thr	Ala .:		٠.	
Ala	Asp	Thr 115		Val	Tyr	Туг	Cys 120		Lys	Ser	Met	I le 125	Thr	Thr	Gly			
Phe	Va 1		: Asp	Ser	Trp	Gly 135		Gly	Ser	Leu								•
(2)	INF	ORM	ATIO	I FOR	SE(	Q ID	NO:	13:		•	×.						**	
j	(		EQUEI	LENG	TH:	436	base	pai	rs				t v *.	- - -	*		:	
			(B) (C) (D)	STRA	NDED	NESS		uble			* •	: a	•			· .~		-
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τ	TCGA	AATG	T CA	ATGN	GTCG	TGT	GTCC'	TGG	AGTG	GTAC	CC T	ACCT	CGAC	A TA	GTAGG	AGA:		60
. A	GAAC	CATC	G TT	GTCG	ATGT	CCA	CAGG	TGA	GGGT	CCAG	GT T	GACG	TCCT	C TC	GCCAG	GTC	1	120
С	AGAA	CACT	C TG	GATC	GGTC	TGG	GACT	CGG	ACTG	GACG	TG G	CACA	GACC	G AA	AAGCA	ATT		180

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GGTCAATACC ACAGGTGACC CACTCTGTCG GTGGA	CCTGC TCCAGAACT	C ACCTAACCTC	240
ACTACACCTC TCCACCTTCG TGTCTGATGT TACGT	CGAAA GTACAGGTC	T GACTTGTAGT	300
GGTTCCTGTT GTCGTTCTTG GTCCACTCGA ATTCT	FGAGTC GTCGCACTG	T CGGCGGCTGT	360
GGCGCCAGAT AATAACACGT TTTAGGTACT ATTGC	CTGGCC GAAGCACTA	C CTGAGGACCC	420
CAGTTCCGAG TGATCA			436
(2) INFORMATION FOR SEQ ID NO: 14:			
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 24 base pairs			
(B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear			.*
(xi) SEQUENCE DESCRIPTION: SEQ ID	) NO: 14:		0
GACATTCAGC TGACCCAGTC TCCA	~		24
(2) INFORMATION FOR SEQ ID NO: 15:			
(i) SEQUENCE CHARACTERISTICS:  (A) LENGTH: 34 base pairs  (B) TYPE: nucleic acid  (C) STRANDEDNESS: single			
(D) TOPOLOGY: linear	•		

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 15:

GATCAAGCTT GACATTCAGC TGACCCAGTC TCCA

- (2) INFORMATION FOR SEQ ID NO: 16:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 37 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 16:

ACTAGTEGAC CATCETECTT TTETGTTTET CTAGGAG

37

- (2) INFORMATION FOR SEQ ID NO: 17:
- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 30 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 17:

		KOON

#### PCT/GB95/02777

GATCAAGCTT	CTCTACAGTT	ACTGAGCACA
	•	

30

#### (2) INFORMATION FOR SEQ ID NO: 18:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 42 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 18:

## CCGATTATAT ATGTCCTCAC TTGCCTTACA GGTGATGGTC AC

42

- (2) INFORMATION FOR SEQ ID NO: 19:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 42 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

### (xi) SEQUENCE DESCRIPTION: SEQ ID NO: 19:

### AGTGAGGACA TATATAATCG GTTAACCTGG TACCAGCAGA AG

42

# (2) INFORMATION FOR SEQ ID NO: 20:

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- (A) LENGTH: 39 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 20:

# AGTITCCAAA CTGGTTGCAC CAGAGATCAG CAGCTTTGG

39

# (2) INFORMATION FOR SEQ ID NO: 21:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 36 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 21:

# GGTGCAACCA GTTTGGAAAC TGGTGTGCCA AGCAGA

36

# (2) INFORMATION FOR SEQ ID NO: 22:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 39 base pairs
  - (B) TYPE: nucleic acid

(C)	STRANDEDNESS:	single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 22:

GTACGGATTA CTCCAATACT GTTGGCAGTA GTAGGTGGC

39

# (2) INFORMATION FOR SEQ ID NO: 23:

### (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 39 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 23:

CAGTATTGGA GTAATCCGTA CACGTTCGGC CAAGGGACC

39

# (2) INFORMATION FOR SEQ ID NO: 24:

### (i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 30 base pairs
- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: single
- (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 24:

GATCAAGCTT CTAACACTCT CCCCTGTTGA

30

- (2) INFORMATION FOR SEQ ID NO: 25:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 30 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 25:

GATCAAGCTT TACAGTTACT CAGCACACAG

- (2) INFORMATION FOR SEQ ID NO: 26:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 30 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 26:

GTGGACACCA TAACTGGTGA AGGTGAAGCC

30

# (2) INFORMATION FOR SEQ ID NO: 27:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 30 base pairs
  - (B) TYPE: nucleic acid
  - (C) STRANDEDNESS: single
  - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 27:

AGTTATGGTG TCCACTGGGT GAGACAGCCA

30

- (2) INFORMATION FOR SEQ ID NO: 28:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 47 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 28:

TTGTAGTCTG TGCTTCCACC TCTCCACATC ACTCCAATCC ACTCAAG

(2)	INFORMATION	FOR	SEQ	ID	NO:	29:
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# (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 51 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 29:

GGAGTCCATC ACGAAGCCGG TCGTTATCAT GGATTTTGCA CAATAATAGA C

51

# (2) INFORMATION FOR SEQ ID NO: 30:

# (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 72 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 30:

AAATCCATGA TAACGACCGG CTTCGTGATG GACTCCTGGG GTCAAGGCTC ACTAGTCACA

72

60

GTCTCCTCAG CC

(2) INFORMATION FOR SEQ ID NO: 31:

	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 36 base pairs	
	(B) TYPE: nucleic acid	
	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
	(D) TOPOLOGI. Tilledi	
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	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 31:	
٠.		
٠.	TAGAGTCCTG AGGGAATTCG GACAGCCGGG AAGGTG	36
	(2) INFORMATION FOR SEQ ID NO: 32:	
· · .		, V
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 39 base pairs	
	(B) TYPE: nucleic acid	
٠. ٠	(C) STRANDEDNESS: single	
	(D) TOPOLOGY: linear	
**		. (1)
		• **
٠.		
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 32:	
	GTTGTCCTTG GTGATGTTCA GTCTGGACAT GAAAGCTGC	3
*	(2) INFORMATION FOR SEQ ID NO: 33:	
	(i) SEQUENCE CHARACTERISTICS:	
	(A) LENGTH: 39 base pairs	,
-	ful remain. 22 page hairs	

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 33:

CTGAACATCA CCAAGGACAA CAGCAAGAAC CAGTTCAGC

39

- (2) INFORMATION FOR SEQ ID NO: 34:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 36 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 34:

ACTGGTTAAC GAAAAGCCAG ACACGGTGCA GGTCAG

- (2) INFORMATION FOR SEQ ID NO: 35:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 36 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID N	0: 35:	**	
GGCTTTTCGT TAACCAGTTA TGGTGTCCAC TGGGTG			36
(2) INFORMATION FOR SEQ ID NO: 36:		e. e	·
(i) SEQUENCE CHARACTERISTICS:			
(A) LENGTH: 36 base pairs	9		* * ÷
(B) TYPE: nucleic acid			
(C) STRANDEDNESS: single			
(D) TOPOLOGY: linear		*	
		* *	
(xi) SEQUENCE DESCRIPTION: SEQ ID N	NO: 36:	*	. 8
AAATTGCCGT TTCGAAGTGT CTACCAGCAT TGTCAC	•		36
(2) INFORMATION FOR SEQ ID-NO: 37:			
(i) SEQUENCE CHARACTERISTICS:			
(A) LENGTH: 36 base pairs		X.	
(B) TYPE: nucleic acid		a 10 -	
(C) STRANDEDNESS: single			
(D) TOPOLOGY: linear		Section 2	

(xi	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:	37

AAATTGCCGT TTCGAATTGT CCTTGGTGAT GTTCAG

36

- (2) INFORMATION FOR SEQ ID NO: 38:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 34 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 38:

TTCGAAACGG CAATTTAGCT TGAGACTCAG CAGC

34

- (2) INFORMATION FOR SEQ ID NO: 39:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 47 base pairs
  - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: single
    - (D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 39:.

GAAGCACAGA CTACAATGCA GCTTTCATGT CCAGAGTGAC AATGCTG

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۱	(4)	INFORMATION	<b>FUK</b>	3EŲ	ΤD	NU:	.40

### (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 28 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 40:

GATCAAGCTT TACAGTTACT CAGCACAG

28

### (2) INFORMATION FOR SEQ ID NO: 41:

### (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 36 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 41:

ACTGGTTAAC GAAAAGCCAG ACACGGTGCA GGTCAG

36

### (2) INFORMATION FOR SEQ ID NO: 42:

(i) SEQUENCE CHARACTERISTICS:

**69** 

(A) LENGTH: 36 base pairs

(B) TYPE: nucleic acid		
(C) STRANDEDNESS: single		
(D) TOPOLOGY: linear		•
	· • • • • • • • • • • • • • • • • • • •	
	**	-
*		* *
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:	42:	
(21) 00(001000		
GCTTTTCGT TAACCAGTTA TGGTGTCCAC TGGGTG	V a	36
gerrirea.		
2) INFORMATION FOR SEQ ID NO: 43:		
2) THE ORBITION TON SEQ SE		*
(i) SEQUENCE CHARACTERISTICS:	*	
(A) LENGTH: 36 base pairs		
(B) TYPE: nucleic acid		,
(C) STRANDEDNESS: single		#10 SL
(D) TOPOLOGY: linear	•	**
(D) TOPOLSON TIMES		
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والمعارية والمراز والمناف والمناف والمناف		
	4	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO	: 43:	
(XI) SEQUENCE SECONDS		
TAGAGTCCTG AGGGAATTCG GACAGCCGGG AAGGTG		36
TAGAGICCI AGGENITO		•
(2) INFORMATION FOR SEQ ID NO: 44:		
(2) IN ORBATZON TON OUT	e de la companya del companya de la companya del companya de la co	,
(i) SEQUENCE CHARACTERISTICS:	•	•
(A) LENGTH: 33 bas pairs		
(B) TYPE: nucleic acid		
(C) STRANDEDNESS: single	*	
(C) SIRANDEDRESS. SINGI		

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(xi)	SEQUENCE	DESCRIPTION:	SEQ	ID	NO:	44:
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## AACCAGGTGA GCTTAAGACT CAGCAGCGTG ACA

33

## (2) INFORMATION FOR SEQ ID NO: 45:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 33 base pairs

(B) TYPE: nucleic acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 45:

### TCTTAAGCTC ACCTGGTTCT TGCTGTTGTC CTT

33

### (2) INFORMATION FOR SEQ ID NO: 46:

## (i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 5 amino acids

(B) TYPE: amino acid

(C) STRANDEDNESS: single

(D) TOPOLOGY: linear

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71

(xi) SEQUENCE DESCRIPTION: SEQ ID NO: 46:

Ser Tyr Gly Val His
1 5

10

#### CLAIMS:

- 1. A monoclonal antibody having donor CDRs of foreign origin and a recipient framework region having a sequence of human or primate origin, wherein the original amino acid residue in position 29 or 78 of the sequence of the recipient framework region of the heavy chain is replaced by a replacement amino acid residue that is the same or similar to that in the corresponding position of the sequence of the corresponding framework region of the heavy chain of the antibody from which the CDRs are derived.
- 2. A monoclonal antibody according to claim 1, wherein the original amino acid residues in both positions 29 and 78 of the sequence of the recipient framework region\_of the heavy chain are replaced by replacement amino acids that are the same or similar to the amino acids in the corresponding positions of the corresponding framework region of the antibody from which the CDRs are derived.
  - 3. A monoclonal antibody according to claim 1 or 2, wherein one or both of the original amino acid residues of the recipient framework region are replaced by a replacement amino acid residues of similar size, hydrophobicity and charge to the amino acids in the corresponding positions of the corresponding framework region of the antibody from which the CDRs are derived.
- 4. A monoclonal antibody according to any of the preceding claims, wherein the original amino acid residues of the recipient framework region are the same or different and are tyrosine, histidine, tryptophan or 2-phenyl-alanine.

- 5. A monoclonal antibody according to claim 4, wherein the replacement amino acid residues are the same or different and are selected from glycine, alanine, valine, serine or leucine.
- 6. A monoclonal antibody according to any of the preceding claims wherein the recipient framework region is from a heavy chain selected from LES-C, T52, Ab44, HIGI and NEW.
- 7. A monoclonal antibody according to any of the preceding claims, wherein the CDRs are of rat, mouse rabbit, or hamster origin.
- 15 8. A monoclonal antibody according to any of the preceding claims, wherein the heavy chain of the antibody from which the CDRs are derived is a murine heavy chain in Kabat groups IB and IIC.
- 20 9. A monoclonal antibody according to any of the preceding claims wherein the antibody binds to CD38.
  - 10. A monoclonal antibody according to claim 9 having a nucleotide sequence as shown in figures 3, 3a and 4.
  - 11. A monoclonal antibody according to any of the preceding claims, wherein the donor CDR is CDRHI.
  - 12. A monoclonal antibody according to claim 11, wherein 30 CDRHI has a sequence of SYGVH.
    - 13. A m thod of producing an antibody according to any of the above claims comprising the st ps of:
  - 35 (i) obtaining the sequence of a donor heavy chain;

- (ii) selecting a recipient human or primate
  framework by best-fit homology method;
- (iii) replacing the amino acid residue in position
  29 or 78 of the sequence of the recipient
  framework region of the heavy chain by an
  amino acid that is the same or similar to that
  in the corresponding position of the sequence
  of the corresponding framework region of the
  antibody from which the CDRs are derived;
  - (iv) grafting donor CDRs into the recipient human framework.
- 15 14. Use of an antibody according to any of the preceding claims for the treatment of cancer and autoimmune diseases.
- 15. Use of an antibody according to claim 9 or 10 for treatment of multiple myeloma, lymphoma and autoimmune diseases such as rheumatoid arthritis.
- 16. Use of an antibody according to any of claims 1 to 12 for the manufacture of a medicament for the treatment 25 of cancer or an autoimmune disease.
  - 17. Use of an antibody according to any of claims 1 to 12 for the manufacture of a medicament for the treatment of multiple myeloma, lymphoma, or rheumatoid arthritis.
  - 18. A pharmaceutical composition comprising an antibody according to any of claims 1 to 12 and a physiologically acceptable diluent or carrier.

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61	TA					TTC			GTA	TAT	ATT	AGC	CAA	TTG	GAC	CAT	AGT	CGT	CIT	TGGT	
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	GG	AAA:	rgc'	rcc	TAG	GCT	CIT	AAT	ATC	TGG	TGC	AAC	CAG	TTT	GGA	AAC	TGG +	GGT	TCC	TTCA	180
121	CC	TIT	ACG	AGG	ATC	CGA	GAA	TTA	TAG	ACC	ACC	TTC	GTC	AAA	CCT	TTG	ACC	CCA	AGG	AAGT	
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		AGA	TGI	TGC	TAC	CCT	ATT!	ACTO	STC	AC	AGT	ATT	GGA(	GTA	ATC	GTA	CAC	GII	CGG	AGGG	300
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Fig.2

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Fig.3a

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	61	AGA		+ TCG'	TTG	rcga	TG	+ TCC	ACA	AGG1	GA(	GG1	rccz	\GG1	TGA	CGT	CCI	CTC	GCC	:AGG	TC	
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	•	GTC	TGI	'GAG	ACC	TAG	CCA	GAC	:CC	TGA(	GCC,	TGA	CCT	ECA(	CG1	rGTC	TGG	+		CGT1	-+	180
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Fig.4

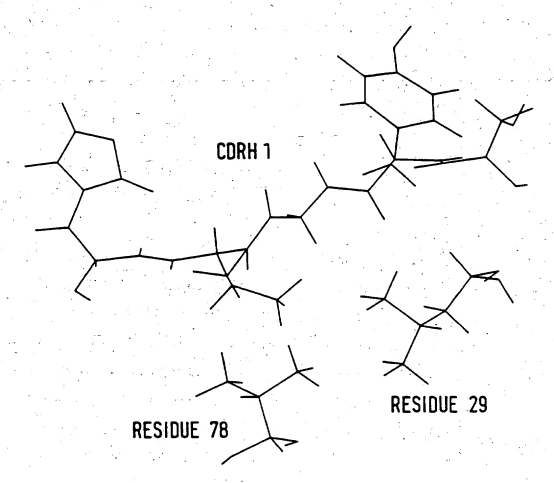


Fig. 5

Fig. 6

RESIDUE 78

# EFFECT OF VARIOUS HEAVY CHAIN FRAMEWORK SUBSTITUTIONS ON RELATIVE BINDING AFFINITY OF ANT-CD38 ANTIBODIES

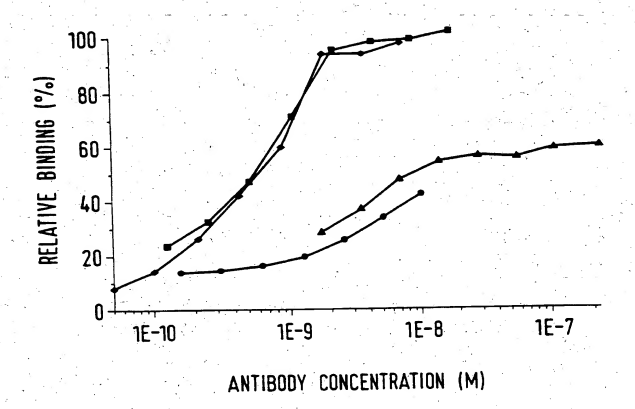
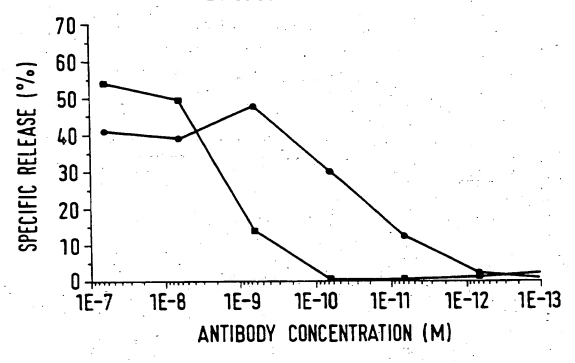


Fig.7

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EFFECT OF VARIOUS HEAVY CHAIN FRAMEWORK SUBSTITUTIONS ON ANTIBODY-DEPENDENT CELLULAR CYTOTOXICITY MEDICATED BY CD38 ANTIBODIES



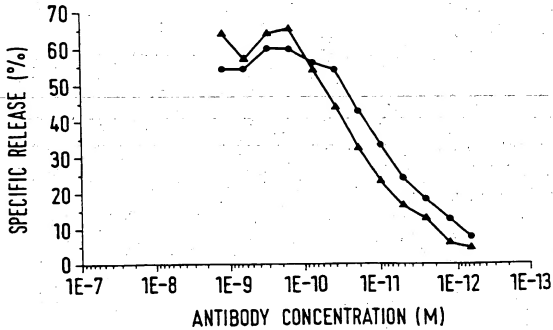


Fig. 8

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	nternational Patent Classification (IPC) or to both national classi	fication and IPC
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PC 6	CO7K A61K	
ocumentation	searched other than minimum documentation to the extent that	such documents are included in the fields searched
lectronic data	a base consulted during the international search (name of data ba	ise and, where practical, search terms used)
	NTS CONSIDERED TO BE RELEVANT	The state of the No.
Category (	Citation of document, with indication, where appropriate, of the	relevant passages Relevant to claim No.
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	see claims	
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X Furt	her documents are listed in the continuation of box C:	X Patent family members are listed in annex.
* Special ca	tegories of cated documents :	"T" later document published after the international filing date or priority date and not in conflict with the application but
'A' docum	nent defining the general state of the art which is not	ated to understand the principle or theory underlying the
consid	document but published on or after the international	"X" document of particular relevance; the claimed invention
ជារកន :	date	cannot be considered novel or cannot be considered to myolve an inventive step when the document is taken alone
which	sent which may throw doubts on priority claim(s) or its cited to establish the publication date of another	"Y" document of particular relevance; the claimed invention
Carego (O. 1	on or other special reason (as specified) ment referring to an oral disclosure, use, exhibition or	document is combined with one or more other such docu- ments, such combination being obvious to a person stalled
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Date of the	e actual completion of the international search	Date of mailing of the international search report
	*	2 3. 04. 96
2	27 March 1996	•
Name and	mailing address of the ISA	Authorized officer
	European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Ripswijk	
1	Tel. (+ 31-70) 340-2040, Tx. 31 651 epo nl.	Nooij, F

Int onal Application No PCT/GB 95/02777

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WO.A.94 13805 (CELLTECH LTD.) 23 June 1994   1,3,4,7,6,13,14,16,18   16,18	C.(Continue		
see page 12, line 19 - page 13, line 5 see example 1 see claims  ( WO,A,91 09967 (CELLTECH LTD.) 11 July 1991 1,3,5,7, 13-18  see table 2 see examples see claims  ( WO,A,94 09136 (KETTOCK LODGE, CAMPUS 2) 28 April 1994 see claims  X JOURNAL OF MOLECULAR BIOLOGY, vol. 224, no. 2, 20 March 1992 LONDON, GB, pages 487-499, XP 000564649 J. FOOTE ET AL. 'Antibody framework residues affecting the conformation of the hypervariable loops.' see abstract see table 2  X PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE USA, vol. 89, no. 10, 15 May 1992 WASHINGTON, DC, USA, pages 4285-4289, XP 000275844 P. CARTER ET AL. 'Humanization of an anti-pla5HER2 antibody for human cancer therapy.' see abstract see table 1  A EP,A,0 481 790 (THE WELLCOME FOUNDATION) 22 April 1992 see claims  THE JOURNAL OF IMMUNOLOGY, vol. 155, no. 2, 15 July 1995 BALTIMORE, MD, USA, pages 925-937, J. ELLIS ET AL. 'Engineered anti-CD38 monoclonal antibodies for immunotherapy of multiple myeloma.'	Category *	Citation of document, with indication, where appropriate, of the relevant passages	Relevant to claim No.
see example 1 see claims  WO,A,91 09967 (CELLTECH LTD.) 11 July 1991  1,3,5,7, 13-18  see table 2 see examples see claims  WO,A,94 09136 (KETTOCK LODGE, CAMPUS 2) 28 April 1994 see claims  X JOURNAL OF MOLECULAR BIOLOGY, vol. 224, no. 2, 20 March 1992 LONDON, GB, pages 487-499, XP 000564649 J. FOOTE ET AL. 'Antibody framework residues affecting the conformation of the hypervariable loops.' see abstract see table 2  X PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE USA, vol. 89, no. 10, 15 May 1992 WASHINGTON, DC, USA, pages 4285-4289, XP 000275844 P. CARTER ET AL. 'Humanization of an anti-p185HER2 antibody for human cancer therapy.' see abstract see table 1  A EP,A,0 481 790 (THE WELLCOME FOUNDATION) 22 April 1992 see Claims  P,X THE JOURNAL OF IMMUNOLOGY, vol. 155, no. 2, 15 July 1995 BALTIMORE, MD, USA, pages 925-937, J. ELLIS ET AL. 'Engineered anti-CD38 monoclonal antibodies for immunotherapy of multiple myeloma.'	X		8,13,14,
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see table 2 see examples see claims  WO,A,94 09136 (KETTOCK LODGE, CAMPUS 2) 28 April 1994 see claims  X JOURNAL OF MOLECULAR BIOLOGY, vol. 224, no. 2, 20 March 1992 LONDON, GB, pages 487-499, XP 000564649 J. FOOTE ET AL. 'Antibody framework residues affecting the conformation of the hypervariable loops.' see abstract see table 2  X PROCEEDINGS OF THE NATIONAL ACADEMY OF SCIENCES OF THE USA, vol. 89, no. 10, 15 May 1992 WASHINGTON, DC, USA, pages 4285-4289, XP 000275844 P. CARTER ET AL. 'Humanization of an anti-pl85HER2 antibody for human cancer therapy.' see abstract see table 1  A EP,A,0 481 790 (THE WELLCOME FOUNDATION) 22 April 1992 see claims  P,X' THE JOURNAL OF IMMUNOLOGY, vol. 155, no. 2, 15 July 1995 BALTIMORE, MD, USA, pages 925-937, J. ELLIS ET AL. 'Engineered anti-CD38 monoclonal antibodies for immunotherapy of multiple myeloma.'	Κ .	WO,A,91 09967 (CELLTECH LTD.) 11 July 1991	
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ternational application No.

## INTERNATIONAL SEARCH REPORT

PCT/GB95/02777

Box	1 Observations where certain claims were found unsearchable (Continuation of item 1 first sheet)
	and the following reasons:
This	international search report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:
. 1	X Claims Nos.: 14,15
յ 1. [	because they relate to subject matter not required to be searched by this Authority, namely:
	Remark: Although claims 14 and 15 are directed to a method of treatment of the human/animal body, the search has been carried out and based on
	the alleged effects of the compound/composition.
2.	Claims Nos.:
	Claims Nos.:  because they relate to parts of the international application that do not comply with the prescribed requirements to such an extent that no meaningful international search can be carried out, specifically:
3.	Claims Nos.:
3.	Claims Nos.:  because they are dependent claims and are not drafted in accordance with the second and third sentences of Rule 6.4(a).
Bo	Il Observations where unity of invention is lacking (Continuation of item 2 of first sheet)
	s International Searching Authority found multiple inventions in this international application, as follows:
Th	s international Searching Additiontly found in-Capita inventions in the international Searching Additiontly found in-Capita inventional in-Capita
}	
1.	As all required additional search fees were timely paid by the applicant, this international search report covers all searchable claims.
2.	As all searchable claims could be searches without effort justifying an additional fee, this Authority did not invite payment
	of any additional fee.
	As only some of the required additional search fees were timely paid by the applicant, this international search report
3.	As only some of the required anditional search rees were drively paid by driving covers only those claims for which fees were paid, specifically claims Nos.:
1	Community this international search report is
4.	No required additional search fees were timely paid by the applicant. Consequently, this international search report is restricted to the invention first mentioned in the claims; it is covered by claims Nos.:
•	
R	emark on Protest  The additional search fees were accompanied by the applicant's protest.
	No protest accompanied the payment of additional search fees.
ı	

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